Evaluation Report

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Fluopyram

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Overview

Registration Decision for Fluopyram

Health Canada's Pest Management Regulatory Agency (PMRA), under the authority of the *Pest Control Products Act* and Regulations, has granted conditional registration for the sale and use of the technical active, Fluopyram Technical Fungicide and end-use products, Luna Privilege containing the technical grade active ingredient fluopyram, Luna Tranquility Fungicide containing the technical grade active ingredients fluopyram and pyrimethanil, and Propulse Fungicide containing the technical grade active ingredients fluopyram and prothioconazole. All three end-use products are used to control several fungal diseases on various horticultural and field crops.

An evaluation of available scientific information found that, under the approved conditions of use, the products have value and do not present an unacceptable risk to human health or the environment.

Although the risks and value have been found acceptable when all risk reduction measures are followed, the applicant must submit additional scientific information as a condition of registration.

This Overview describes the key points of the evaluation, while the Science Evaluation provides detailed technical information on the human health, environmental and value assessments of fluopyram in Fluopyram Technical Fungicide, Luna Privilege, Luna Tranquility Fungicide and Propulse Fungicide.

What Does Health Canada Consider When Making a Registration Decision?

The key objective of the *Pest Control Products Act* is to prevent unacceptable risks to people and the environment from the use of pest control products. Health or environmental risk is considered acceptable¹ if there is reasonable certainty that no harm to human health, future generations or the environment will result from use or exposure to the product under its proposed conditions of registration. The Act also requires that products have value² when used according to the label directions. Conditions of registration may include special precautionary measures on the product label to further reduce risk.

[&]quot;Acceptable risks" as defined by subsection 2(2) of the Pest Control Products Act.

[&]quot;Value" as defined by subsection 2(1) of the *Pest Control Products Act*: "the product's actual or potential contribution to pest management, taking into account its conditions or proposed conditions of registration, and includes the product's (a) efficacy; (b) effect on host organisms in connection with which it is intended to be used; and (c) health, safety and environmental benefits and social and economic impact."

To reach its decisions, the PMRA applies modern, rigorous risk-assessment methods and policies. These methods consider the unique characteristics of sensitive subpopulations in humans (for example, children) as well as organisms in the environment (for example, those most sensitive to environmental contaminants). These methods and policies also consider the nature of the effects observed and the uncertainties when predicting the impact of pesticides. For more information on how the PMRA regulates pesticides, the assessment process and risk-reduction programs, please visit the Pesticides and Pest Management portion of Health Canada's website at healthcanada.gc.ca/pmra.

What Is Fluopyram?

Fluopyram is a new systemic fungicidal compound present as the lone active ingredient in the new end-use product Luna Privilege. It is also present as one of the two active ingredients in two new pre-mix end-use products, Luna Tranquility Fungicide and Propulse Fungicide. The compound is used as a broad-spectrum fungicide applied as a foliar spray or through drip irrigation systems on various horticultural and field crops. It acts on pathogen cells by inhibiting their normal respiration process.

Health Considerations

Can Approved Uses of Fluopyram Affect Human Health?

Products containing fluopyram are unlikely to affect your health when used according to label directions.

Exposure to fluopyram may occur through the diet (food and water), when handling and applying the product or when entering treated sites. When assessing health risks, two key factors are considered: the levels where no health effects occur and the levels to which people may be exposed. The dose levels used to assess risks are established to protect the most sensitive human population (for example, children and nursing mothers). Only uses for which the exposure is well below levels that cause no effects in animal testing are considered acceptable for registration.

Toxicology studies in laboratory animals describe potential health effects from varying levels of exposure to a chemical and identify the dose where no effects are observed. The health effects noted in animals occur at doses more than 100-times higher (and often much higher) than levels to which humans are normally exposed when pesticide products are used according to label directions.

In laboratory animals, the acute toxicity of fluopyram was low via the oral, dermal and inhalation routes of exposure. Fluopyram was minimally irritating to the eyes and non-irritating to the skin and did not cause an allergic skin reaction.

The acute toxicity of the end-use product Luna Privilege was low via the oral, dermal and inhalation routes of exposure. It was minimally irritating to the eyes and non-irritating to the skin and did not cause an allergic skin reaction. Both end-use products Luna Tranquility Fungicide and Propulse Fungicide were of low acute toxicity via the oral, dermal and inhalation routes of exposure. They were non-irritating to the eyes and skin and did not cause allergic skin reactions.

Health effects in animals given repeated doses of fluopyram included changes in the liver, thyroid and kidneys. Fluopyram did not cause birth defects in animals and there were no effects on the ability to reproduce. When fluopyram was given to pregnant or nursing animals, effects on the developing fetus and juvenile animal (reduced pup and litter weights, body size, spleen and thymus weights, and/or slightly delayed sexual development) were observed at doses that were toxic to the mother, indicating that the young do not appear to be more sensitive to fluopyram than the adult animal. Fluopyram did not selectively target the nervous system, however, temporary non-specific functional effects (decreased motor and locomotor activity) were observed, possibly related to the nervous system. There was no evidence to suggest that fluopyram damaged genetic material. Fluopyram did, however, cause thyroid tumours in mice and liver tumours in rats.

The risk assessment protects against the effects of fluopyram by ensuring that the level of human exposure is well below the lowest dose at which these effects occurred in animal tests.

Residues in Water and Food

Dietary risks from food and water are not of concern.

Aggregate dietary intake estimates (food plus water) revealed that the general population and infants less than one year old, the subpopulation which would ingest the most fluopyram relative to body weight, are expected to be exposed to less than 64% of the acceptable daily intake. Based on these estimates, the chronic dietary risk from fluopyram is not of concern for all population subgroups. The lifetime cancer risk from the use of fluopyram on various crops is considered acceptable, based on a limited three-year application period.

Acute dietary (food and water) estimates for the general population and all population subgroups were less than 10% of the acute reference dose, and are not of health concern. The highest exposed subpopulation was children 1-2 years old.

The Food and Drugs Act prohibits the sale of adulterated food, that is, food containing a pesticide residue that exceeds the established maximum residue limit (MRL). Pesticide MRLs are established for Food and Drugs Act purposes through the evaluation of scientific data under the Pest Control Products Act. Food containing a pesticide residue that does not exceed the established MRL does not pose an unacceptable health risk.

Residue trials conducted throughout Canada and the United States using fluopyram on potatoes, sugar beets, dry beans and dry peas, watermelon, apples, cherries, strawberries, grapes, almonds, pecans, peanuts, soybeans, wheat, sorghum, corn (field and sweet), canola and cottonseed, as well as trials conducted in Latin America using fluopyram on bananas are acceptable. The MRLs for this active ingredient can be found in the Science Evaluation of this Evaluation Report.

Risks in Residential and Other Non-Occupational Environments

Residential risks are not of concern when products containing fluopyram are used according to the label directions.

Occupational Risks from Handling Luna Privilege, Propulse Fungicide and Luna Tranquility Fungicide

Occupational risks are not of concern when products containing fluopyram are used according to the label directions, which include protective measures.

Farmers and custom applicators who mix, load or apply fluopyram as well as field workers reentering freshly treated fields can come in direct contact with fluopyram residues on the skin. Therefore, the labels specify that anyone mixing/loading and applying products containing fluopyram must wear a long-sleeved shirt, long pants, shoes plus socks, and chemical resistant gloves. The label also requires that workers do not enter treated fields for 12 hours after application. Taking into consideration these label statements, the number of applications and the expectation of the exposure period for handlers and workers, risks to these individuals are not a concern.

For bystanders, exposure is expected to be much less than that for workers and is considered negligible. Therefore, health risks to bystanders are not of concern.

Environmental Considerations

What Happens When Fluopyram is Introduced into the Environment?

When fluopyram is applied as a fungicide on field crops, some of it finds its way into soil and water. In soils, it is persistent and has a potential for long-term accumulation and residue carry over to the following crop season. Fluopyram is stable to hydrolysis, photolysis, aerobic and anaerobic biotransformation in soils and does not form major transformation products in soils under Canadian field use conditions. Fluopyram is moderately mobile in soils and has a potential to leach and contaminate the groundwater depending on the soil type and location. None of the minor transformation products, however, have a potential to leach and contaminate groundwater. Fluopyram has a low potential for bioconcentration/bioaccumulation in organisms.

In the aquatic environment, fluopyram is persistent under aerobic and anaerobic conditions and partitions significantly from water to sediment. It does not form any major transformation products in water or sediment phases. Photolysis is not an important route of transformation in the aquatic environment. Several minor transformation products were detected due to photolysis under laboratory conditions in natural water of which one was identified as fluopyram-lactam.

Fluopyram has a low potential for volatilization and, therefore, not expected to result in long range transport in the atmosphere.

Fluopyram presents a negligible risk to soil organisms, bees, beneficial arthropods, freshwater and marine fish, invertebrates, algae and aquatic plants. Fluopyram, however, may pose a risk to non-target terrestrial plants from spray drift (Luna Privilege only), and to amphibians due to runoff and spray drift. In order to minimize the potential risk, no-spray buffer zones between the treated area and downwind sensitive terrestrial and aquatic habitats are required. A bird toxicity label statement is also required as a precaution.

Value Considerations

What Is the Value of Luna Privilege, Luna Tranquility Fungicide and Propulse Fungicide?

Luna Privilege, Luna Tranquility Fungicide and Propulse Fungicide are fungicides effective in the control of major economic diseases of various horticultural and field crops.

Luna Privilege, Luna Tranquility Fungicide and Propulse Fungicide provide effective solutions for the management of major economic diseases such as powdery mildew, moulds, blights and other foliar diseases on a range of crops including potato, dry bean, chickpeas, lentils, apple, cherry, wine grape, strawberry, peanut, watermelon and almond. The combinations of different modes of action in Luna Tranquility Fungicide and Propulse Fungicide are of value in reducing the risk of resistance development and by increasing the spectrum of disease protection.

Measures to Minimize Risk

Labels of registered pesticide products include specific instructions for use. Directions include risk-reduction measures to protect human and environmental health. These directions must be followed by law.

The key risk-reduction measures on the label of Luna Privilege, Luna Tranquility Fungicide, and Propulse Fungicide to address the potential risks identified in this assessment are as follows.

Key Risk-Reduction Measures

Human Health

Because there is a concern with users coming into direct contact with fluopyram residues on the skin or through inhalation of spray mist, anyone mixing, loading and applying products containing fluopyram must wear a long-sleeved shirt, long pants, shoes plus socks, and chemical resistant gloves. In addition, standard label statements to protect against drift during application were added to the label.

Environment

Based on the risk identified to off-target sensitive habitats, buffer zones of 1 to 15 m are required to protect amphibians and terrestrial habitats. In addition, standard label statements were added to the labels to protect wild birds, aquatic organisms and non-target terrestrial plants.

What Additional Scientific Information Is Being Requested?

Although the risks and value have been found acceptable when all risk-reduction measures are followed, the applicant must submit additional scientific information as a condition of registration. More details are presented in the Science Evaluation section of this Evaluation Report or in the Section 12 Notice associated with these conditional registrations. The applicant must submit the following information within the time frames indicated.

Human Health

- Short-term mode of action studies to address the observed tumours. The goal of these studies is to further clarify the two proposed cancer modes of action.
- Inter-Laboratory Analytical Methodology Validation An independent laboratory validation of Method GM-001-P07-01 for the determination of fluopyram residues in plant matrices is required to fulfill the data requirement for an acceptable enforcement method in plant matrices.
- Field Accumulation Studies A full set of field rotational crop data are required for canola, soybean and cereals (wheat, barley and corn, both field and sweet).

Value

- One field trial to confirm the efficacy of Luna Privilege against powdery mildew on standard sized cherry trees.
- One field trial to confirm efficacy of Luna Privilege against late leaf spot on peanuts.

Other Information

As these conditional registrations relate to a decision on which the public must be consulted,³ the PMRA will publish a consultation document when there is a proposed decision on applications to convert the conditional registrations to full registrations or on applications to renew the conditional registrations, whichever occurs first.

The test data cited in this Evaluation Report (that is, the test data relevant in supporting the registration decision) will be made available for public inspection when the decision is made to convert the conditional registrations to full registrations or to renew the conditional registrations (following public consultation). If more information is required, please contact the PMRA's Pest Management Information Service by phone (1-800-267-6315) or by e-mail (pmra.infoserv@hc-sc.gc.ca).

As per subsection 28(1) of the Pest Control Products Act.

Science Evaluation

Fluopyram

1.0 The Active Ingredient, Its Properties and Uses

1.1 Identity of the Active Ingredient

Active substance Fluopyram Function Fungicide

Chemical name

1. International Union of Pure N-{2-[3-chloro-5-(trifluoromethyl)-2-pyridyl]ethyl}and Applied Chemistry α,α,α-trifluoro-o-toluamide
(IUPAC)

2. Chemical Abstracts Service Benzamide, N-[2-[3-chloro-5-(trifluoromethyl)-2-(CAS) pyridinyl]ethyl]-2-(trifluoromethyl)-

CAS number 658066-35-4

Molecular formula C₁₆H₁₁ClF₆N₂O

Molecular weight 396.72 g/mol

Structural formula F₃C Cl CF₃

Purity of the active ingredient 98.6%

1.2 Physical and Chemical Properties of the Active Ingredients and End-use Product

Technical Product—Fluopyram Technical Fungicide

Property	Result	
Colour and physical state	White powder	
Odour	No noticeable odour	
Melting range	118°C	
Boiling point or range	319°C (correlated range) under decomposition	
Relative Density	1.53	
Vapour pressure at 20°C	$1.2 \times 10^{-6} \text{ Pa } (20^{\circ}\text{C})$	
Henry's law constant at 20°C	$2.98 \times 10^{-5} \text{ Pa} \times \text{m}^3 \times \text{mol}^{-1}$	

Property	Result	
Ultraviolet (UV)-visible spectrum	Acetonitrile:	
•	$\lambda_{\max}[nm]$ $\epsilon [L mol^{-1} cm^{-1}]$	
	216 14877	
	270 4332.18	
	Acetonitrile pH = 2:	
	$\lambda_{\text{max}}[\text{nm}]$ $\varepsilon [\text{L mol}^{-1} \text{cm}^{-1}]$	
	208 16570.99	
	270 4399.62	
	Acetonitrile pH = 10:	
	$\lambda_{\text{max}}[\text{nm}]$ $\epsilon [\text{L mol}^{-1} \text{cm}^{-1}]$	
	208 16892.34	
	270 4383.76	
	Water	
	$\lambda_{\max}[nm]$ $\epsilon [L mol^{-1} cm^{-1}]$	
	270 4577.053	
Solubility in water at 20°C	16 mg/L (distilled water)	
	15 mg/L (pH 4)	
	16 mg/L (pH 7)	
	15 mg/L (pH 9)	
Solubility in organic solvents at 20°C	Solvent Solubility (g/L)	
	acetone >250	
	dichlorethane >250	
	dimethyl sulfoxide >250	
	ethyl acetate >250	
	n-heptane 0.66	
	methanol >250	
	toluene 62.2	
n-Octanol-water partition coefficient (Ko	ow) $\log K_{ow} = 3.3$ at 20° C	
Dissociation constant (pK _a)	No dissociation observed between pH 2 and 12	
Stability	Stable in presence of metals (iron and aluminum) as	
(temperature, metal)	when stored for two weeks at 54°C in presence of	
	metals and metal ions.	

End-use Products-Fluopyram

Property		Result		
	Luna Privilege	Luna Tranquility Fungicide	Propulse Fungicide	
Colour	Beige	Off-white	Off-white	
Odour	Chemical odour	Wine-like odour	Mild sweet odour	
Physical state	Liquid	Liquid	Liquid	
Formulation type	Suspension	Suspension	Suspension	
Guarantee	Fluopyram 500 g/L	Fluopyram 125 g/L Pyrimethanil 375 g/L	Fluopyram 200 g/L Prothioconazole 200 g/L	
Container material and description	HDPE bottle/canister, 0.25 – 10 L, or canister/IBC such as 1000 L	HDPE containers, 1 to 200 L	HDPE containers, 1 to 200 L	
Density	1.205 g/mL	1.11 g/mL	1.15 g/mL	
pH of 1% dispersion in water	6.5	7.2	5.0	
Oxidizing or reducing action	None	None	None	
Storage stability	Stable over 12 months in HDPE packaging at ambient temperature.	Stable when stored for 12 months at ambient temperature in commercial packaging	Stable when stored for 12 months at ambient temperature in commercial packaging	
Corrosion characteristics	Not corrosive	Not corrosive	Not corrosive	
Explodability	Not explosive	Not explosive	Not explosive	

1.3 Directions for Use

Luna Privilege, Luna Tranquility Fungicide and Propulse Fungicide are used for the control of powdery mildew, moulds, blights and other foliar diseases on various field and horticultural crops. The products are intended for foliar applications on all crops with the exception of strawberry where applications via chemigation are indicated for Luna Privilege. The application rate ranges for Luna Privilege, Luna Tranquility Fungicide, and Propulse Fungicide, are 150-500 mL/ha, 600-1200 mL/ha and 500-750 mL/ha, respectively.

1.4 Mode of Action

Fluopyram is a new broad-spectrum systemic active ingredient classified as a group 7 fungicide (succinate dehydrogenase inhibitor) by the Fungicide Resistance Action Committee. Fluopyram interferes with the normal respiration process in the cells of pathogenic fungal cells. Fluopyram shows systemic and preventative activity against the ascomycetes, a group of fungi that includes many economically important crop pathogens.

2.0 Methods of Analysis

2.1 Methods for Analysis of the Active Ingredient

The methods provided for the analysis of the active ingredient and the impurities in Fluopyram Technical Fungicide have been validated and assessed to be acceptable for the determinations.

2.2 Method for Formulation Analysis

The methods provided for the analysis of the active ingredients in the formulations have been validated and assessed to be acceptable for use as enforcement analytical methods.

2.3 Methods for Residue Analysis

High-performance liquid chromatography methods with tandem mass spectrometry (HPLC-MS/MS) were developed and proposed for data generation and enforcement purposes. These methods fulfilled the requirements with regards to selectivity, accuracy and precision at the respective method limit of quantitation. Acceptable recoveries (70-110%) were obtained in environmental media. Methods for residue analysis are summarized in Appendix I, Table 1.

HPLC-MS/MS methods developed and proposed for data generation and enforcement purposes in plant and animal commodities fulfilled the requirements with regards to specificity, accuracy and precision at the respective limits of quantitation of the methods. Acceptable recoveries (70-120%) were obtained in plant and animal matrices. The proposed enforcement method for animal commodities was successfully validated in several animal matrices by an independent laboratory. Adequate extraction efficiencies were demonstrated using radiolabelled samples of several crop matrices and livestock tissues analyzed with the respective enforcement methods Appendix I, Table 1.

3.0 Impact on Human and Animal Health

3.1 Toxicology Summary

Fluopyram is a broad spectrum pyridylethylamide fungicide. A detailed review of the toxicological database for fluopyram was conducted. The database consists of the full array of toxicity studies currently required for hazard assessment purposes. The database also includes neurotoxicity and cancer mode of action (MOA) studies. In addition, an acute oral toxicity, a 28 day dietary toxicity and three genotoxicity studies were provided for a plant/soil metabolite. The studies were carried out in accordance with currently accepted international testing protocols and Good Laboratory Practices. The scientific quality of the data is high and the database is considered adequate to define the majority of the toxic effects that may result from exposure to fluopyram. However, additional information is being developed to further elucidate the cancer modes of action.

The absorption, distribution, metabolism and excretion characteristics of single and multiple radiolabelled doses, were evaluated in rats. Orally administered fluopyram was rapidly and extensively absorbed. Time to maximal tissue concentration varied from 0.8 to 15 hours depending on the placement of radiolabel and the dosing regime. The systemic exposure was proportional to the dose and slightly higher in females compared to males. Absorbed fluopyram was widely distributed, with the concentrations in the plasma being exceeded by the maximal levels in each of the following organs and tissues: the liver, kidneys, and Harderian gland in all test groups as well as the carcass, red blood cells, ovaries, thyroid and adrenals in some groups. Excretion of fluopyram was rapid and dose-independent. Fluopyram was eliminated predominantly via the bile, with appreciable amounts also excreted in the urine. After cessation of dosing, organ and tissue concentrations of radioactivity decreased rapidly. There was 0.3-6% of administered dose remaining in the carcass at 168 hours, depending on the radiolabel position, so the potential for accumulation cannot be ruled out. Fecal elimination was essentially complete within 96 hours. Elimination of fluopyram via respired volatiles and CO2 was negligible. The initial elimination half-life ranged from 3.9 to 16.2 hours depending on the radiolabel position and dose level. The terminal elimination half-life increased to a range of 23.6 to 72.9 hours. There were no significant sex- or dose-related differences in the tissue distribution and retention or in the extent or route of elimination.

Fluopyram was extensively metabolized, with the ethyl linking group of the parent as the preferred site for metabolism, resulting in 7-hydroxy and 8-hydroxy metabolites. Further oxidation resulted in an -enol metabolite, which was further conjugated to glucuronic acid. Hydroxylation of the phenyl ring resulted in -phenol and 7-OH-phenol metabolites. Elimination of water from compounds hydroxylated in the ethylene bridge resulted in fluopyram-Z-olefine and E-olefine metabolites (E- and Z-olefine can isomerize into each other). As the double bond of olefine may be a target for epoxidation and a dihydroxy-metabolite (which could result from hydrolysis of an epoxide by epoxide hydrolase) was observed, the olefine was considered to be of potential toxicological significance. All of the hydroxylated metabolites were conjugated primarily to glucuronic acid and to a lesser extent with sulfate. The cleavage of the molecule yielded label-specific metabolites (-benzamide; -pyridyl-acetic acid, -ethyl-diol, -pyridyl carboxylic acid) that represented the most abundant metabolites. These metabolites were further metabolized via oxidation, hydroxylation and conjugation. The phenyl ring moiety was also conjugated with glutathione followed by further degradation to 7-OH-methyl-sulfone, -BA-methyl-sulfone (phenyl label only).

There were sex differences in the quantity of metabolites generated. Fluopyram-7-hydroxy and 7-OH-phenol metabolites were higher in males than females. Females showed higher amounts of 8-hydroxy and -benzamide than males. Low dose females excreted more of phenyl specific -benzamide and -benzoic acid than males. Females treated with the pyridyl label excreted more -pyridyl-acetic acid than males, while males excreted more -ethyl-diol metabolites than females. Parent compound accounted for 0.4/1.9% 3/\Q of the administered dose for the single oral low dose group and 10.5/16.7% 3/\Q of the administered dose for the single oral high dose group. There were no significant differences in metabolism between the doses, or between single and repeat dosing.

The acute toxicity of the active ingredient fluopyram and its three end-use products was low via the oral, dermal and inhalation routes in rats. All four products were non-irritating to minimally irritating to the eyes and non-irritating to the skin of rabbits. None of the products were skin sensitizers in either guinea pigs or mice.

In the short-term oral studies, the liver was the main target organ in mice, rats and dogs. Hepatotoxicity manifested as increased liver weight, liver enlargement, darkening, necrosis and centrilobular and mid-zonal hepatocellular hypertrophy, as well as alterations in clinical chemistry (elevated plasma/serum levels of liver enzymes, cholesterol and/or phospholipids, and triglycerides with decreased albumin). The rat was the most sensitive species following short-term oral dosing. The liver toxicity between mice, rats and dogs was similar with the exceptions of dark livers in the rodents only, and increased cholesterol and hepatocellular macrovacuolation in rats only. Mice and dogs had hepatocellular necrosis, which was not observed in rats. In several studies, effects on the liver at lower doses were mild and considered to be non-adverse, reflecting an adaptive response of the liver rather than overt hepatotoxicity. The spectrum of liver effects and the doses eliciting hepatotoxicity worsened significantly with the duration of dosing (short-term to chronic). At higher doses in mice, decreased pigment and increased vacuolation of the adrenals were noted. For rats, higher dose levels resulted in decreased body weight, increased thyroid hormone levels, vacuolation in the adrenals, pale or dark kidneys, kidneys with cysts or cellular debris, follicular cell hypertrophy of the thyroid, increased thyroid weight, and decreased fore- and hindlimb grip strength. High doses in the dogs resulted in decreased body and thymus weight.

Dermal dosing of rats for 28 days resulted in increased prothrombin time, cholesterol, liver weight and minimal hepatocellular hypertrophy at 1000 mg/kg bw/day, the highest dose tested.

The liver, kidneys, and thyroid were the primary target organs in the mouse and rat with chronic oral dosing. With long-term dosing in mice, the thyroid exhibited increased incidences and severity of follicular cell hyperplasia. Liver enlargement and a variety of histopathological effects were also more frequently observed at doses lower than in the short-term study. At the highest dose tested, mouse body weights were decreased, kidney weights were slightly decreased and the incidence and severity of several renal histopathological effects were significantly increased. In rats, the same liver effects seen in the short-term studies were repeated at similar or lower dose levels. Additionally, altered hepatocyte foci and hepatocellular necrosis were identified following 12 and 24 months of treatment. In rat thyroids, the incidence and severity of follicular cell hypertrophy, hyperplasia and colloid alteration were increased. After 24 months the kidneys of male rats exhibited increased incidences and/or severities of chronic progressive nephropathy, tubular hyperplasia, hypertrophy or dilatation, and golden brown pigment. In rats, the eyes were also a target organ with corneal opacity and edema, opacity of the lens and small retinal vessels seen at relatively low dose levels. At the highest two doses tested in the 12- and 24-month rat studies, there were additional generalized findings such as decreased body weight, prostration, pallor, wasted appearance and hair loss.

Fluopyram was tested for in vitro and in vivo genotoxicity in a range of assays. Based on the negative results obtained in a battery of genotoxicity studies, fluopyram is considered unlikely to be genotoxic.

Tumours were observed in the mouse and the rat in the dietary oncogenicity studies. The dosing was considered adequate in these studies. Male mice had thyroid follicular cell adenomas while female rats had liver adenomas and carcinomas. These tumours are considered uncommon in the respective species/sex. The proposed MOA for the thyroid adenomas was chronic perturbation of

thyroid hormone homeostasis. In liver, the proposed MOA was phenobarbital-like liver proliferation. Cancer MOA studies were conducted to examine liver and thyroid effects in the rat and mouse. These studies in mice showed that fluopyram increased T4 elimination, but did not affect thyroid hormone synthesis. Fluopyram also up-regulated sulfotransferase and UDP glucuronosyltransferase transcripts in the liver. These transcripts are known to encode enzymes that inactivate T3 and T4. Additionally, P450, EROD, PROD, and BROD enzymes were increased in fluopyram treated mice. While the evidence was generally supportive for the thyroid tumour MOA, there are data gaps in terms of dose and time concordance between the MOA data and the tumourigenic dose levels. In female rats, hepatocellular hypertrophy and liver cell proliferation were associated with the induction of xenobiotic metabolizing enzymes. Again, while the evidence was generally supportive for the liver tumour MOA, there were data gaps. Overall, when the results from all of the MOA studies in mice and rats are considered, there was insufficient evidence to conclude that the oncogenic effects in the thyroid and liver were specific consequences of chronically perturbed thyroid hormone homeostasis and chronically induced liver metabolizing enzymes. A linear low dose extrapolation (Q₁*) approach was used for the cancer risk assessment in the absence of a sufficient weight of evidence to support a proposed threshold-based MOA.

No effects on reproduction were noted in a multigeneration reproduction study in the rat. There was a decrease in offspring body weight during early lactation in both generations at the highest dose tested. Also, at this dose, there were decreases in thymus and spleen weights, with no histopathological correlates. Effects were also observed in parental animals at the high dose and included decreased body weight and body weight gain, increased cholesterol and white blood cell counts, increased liver weight with centrilobular hepatocellular hypertrophy, increased kidney weight with nephropathy and lymphocytic infiltration, decreased spleen weight in the absence of histopathological changes, increased vacuolization in the adrenals and macrophages in the lungs. There was no evidence of sensitivity of the young.

In the oral developmental toxicity study in rats, the only fetal effects noted were decreased fetal weight, thymic remnants and four different skeletal variations at the highest dose tested. The dams exhibited decreased body weight gain and food consumption along with increased liver weights and centrilobular hepatocellular hypertrophy starting at the mid dose, plus decreased body weights and visibly enlarged livers at the high dose. In rabbits, fetal weights were decreased and the number of runts was increased at the highest dose tested. The does at that dose level had decreased body weight, body weight gains and food consumption. Fluopyram is not considered teratogenic and it induced fetal toxicity only in the presence of maternal toxicity.

In an acute oral neurotoxicity study in rats, females in all dose groups exhibited decreased session motor and locomotor activity on the day of testing. Males were similarly affected starting at the mid dose. A supplemental study with females at lower doses was able to determine a no observed adverse effect level (NOAEL) for those effects. In the short-term oral neurotoxicity test in rats, there was no evidence of neurotoxicity following dietary administration of fluopyram. Noted effects matched those in the main toxicity studies, namely, decreased food consumption and increased liver, kidney and thyroid weight.

A pyridyl-carboxylic acid metabolite of fluopyram was tested in an acute oral toxicity study and a short-term toxicity study, both in rats. In the acute study, the LD_{50} was greater than 2000 mg/kg bw with piloerection observed following the day of dosing at 500 mg/kg bw. The short-term study resulted in decreased body weight gain and food consumption. The metabolite was less toxic than parent fluopyram in the studies provided.

Results of the toxicology studies conducted on laboratory animals with fluopyram and its associated end-use products are summarized in Appendix I, Tables 2–5. The toxicology endpoints for use in the human health risk assessment are summarized in Appendix I, Table 6.

Incident Reports

Since 26 April 2007, registrants have been required by law to report incidents, including adverse effects to health and the environment, to the PMRA within a set time frame. Information on the reporting of incidents can be found on the Pesticides and Pest Management portion of Health Canada's website at healthcanada.gc.ca/pesticideincident. Incident reports from Canada and the United States were searched for fluopyram and any additional information submitted by the applicant during the review process was considered. As of 13 March 2012, there were no health-related incident reports for this active in the PMRA Incident Reporting database.

3.1.1 Pest Control Products Act Hazard Characterization

For assessing risks from potential residues in food or from products used in or around homes or schools, the *Pest Control Products Act* requires the application of an additional 10-fold factor to threshold effects to take into account completeness of the data with respect to the exposure of, and toxicity to, infants and children, and potential prenatal and postnatal toxicity. A different factor may be determined to be appropriate on the basis of reliable scientific data.

With respect to the completeness of the toxicity database as it pertains to the toxicity to infants and children, extensive data were available for fluopyram. The database contains the full complement of required studies including developmental toxicity studies in rats and rabbits and a reproductive toxicity study in rats.

With respect to potential prenatal and postnatal toxicity, there was no indication of increased susceptibility of fetuses or offspring compared to parental animals in the reproductive toxicity and prenatal developmental toxicity studies. In the 2-generation rat reproductive toxicity study, adverse effects on offspring body size and weight only occurred in the presence of maternal toxicity (liver, adrenal, blood and bodyweight effects). Maternal toxicity (bodyweight effects in both species and liver effects in rats) in the oral developmental toxicity studies in rats and rabbits also tempered concern for the decreased fetal weight in both species, the skeletal variations in rats and the number of runts in rabbits. Fluopyram was not considered teratogenic.

Overall, endpoints in the young were well characterized and not considered serious in nature. The *Pest Control Products Act* factor was reduced to 1-fold. The endpoints selected for risk assessment were protective of the effects noted in the rat and rabbit reproduction and developmental toxicity studies.

3.2 Acute Reference Dose (ARfD)

General Population

To estimate acute dietary risk (one day), the acute oral neurotoxicity study in rats with a NOAEL of 50 mg/kg bw was selected for risk assessment. At the lowest observed adverse effect level (LOAEL) of 100 mg/kg bw, session motor and locomotor activities were decreased in females. These effects were the result of a single exposure and are therefore relevant to an acute risk assessment. Standard uncertainty factors of 10-fold for interspecies extrapolation and 10-fold for intraspecies variability were applied. As discussed in the *Pest Control Products Act* Hazard Characterization section, the *Pest Control Products Act* factor was reduced to 1-fold. The composite assessment factor (CAF) is 100.

The ARfD is calculated according to the following formula:

ARfD (gen. pop.) =
$$\frac{\text{NOAEL}}{\text{CAF}} = \frac{50 \text{ mg/kg bw}}{100} = 0.5 \text{ mg/kg bw}$$

3.3 Acceptable Daily Intake (ADI)

To estimate dietary risk from repeat exposure, the 24-month oral chronic toxicity/oncogenicity study in rats with a NOAEL of 1.2 mg/kg bw/day was selected for risk assessment. At the LOAEL of 6.0 mg/kg bw/day, increases in liver hypertrophy, kidney weight and histopathology, cellular casts in urine, thyroid hypertrophy and colloid alteration and ocular toxicity were all observed. This study provides the lowest NOAEL in the database. Standard uncertainty factors of 10-fold for interspecies extrapolation and 10-fold for intraspecies variability were applied. As discussed in the *Pest Control Products Act* Hazard Characterization section, the *Pest Control Products Act* factor was reduced to 1-fold. The composite assessment factor (CAF) is 100.

The ADI is calculated according to the following formula:

$$ADI = \frac{NOAEL}{CAF} = \frac{1.2 \text{ mg/kg bw/day}}{100} = 0.012 \text{ mg/kg bw/day}$$

Cancer Assessment

Fluopyram showed evidence of oncogenicity in both the rat and the mouse. There was some evidence supporting a threshold-based mechanism to the tumors (thyroid tumours in mice and liver tumours in rats), however, further data are required to establish MOAs. In the interim, a linear low dose extrapolation (Q_1^*) was used for risk assessment, but is considered to be conservative. The Q_1^* was set at 1.72×10^{-2} (mg/kg bw/day)⁻¹.

3.4 Occupational and Residential Risk Assessment

3.4.1 Toxicological Endpoints

Occupational and residential exposure to fluopyram is characterized as short- to long-term and is predominantly by the dermal and inhalation routes.

Short- and Intermediate-term Dermal Exposure

For short- and intermediate-term dermal risk assessment, the short-term dermal toxicity study in rats was selected. At the dose of 1000 mg/kg bw/day, clinical chemistry effects and liver toxicity were evident. A NOAEL of 300 mg/kg bw/day was established.

The target MOE selected for this endpoint is 100. Ten-fold factors were applied each for interspecies extrapolation and intraspecies variability. This MOE is considered to be protective of all adults including pregnant and lactating women and their unborn children, as well as nursing infants and children of exposed female workers.

Short- and Intermediate-term Inhalation Exposure

For short- and intermediate-term exposure via the inhalation route, the 90-day oral toxicity study in rats was selected for risk assessment. A NOAEL of 12.5 mg/kg bw/day was established based on decreased food consumption, liver and kidney toxicity and clinical chemistry alterations at 60.5 mg/kg bw/day. This study provides the lowest short- to intermediate-term toxicity NOAEL in the database. A short-term inhalation study was not available.

The target MOE for these scenarios is 100, which includes uncertainty factors of 10-fold for interspecies extrapolation and 10-fold for intraspecies variability. This study and target MOE are considered to be protective of all populations, including nursing infants and the unborn children of exposed female workers.

Long-term Dermal and Inhalation Exposure

For long-term dermal and inhalation risk assessment, the 24-month oral chronic toxicity study in rats with a NOAEL of 1.2 mg/kg bw/day was selected for risk assessment. At the LOAEL of 6.0 mg/kg bw/day, increases in liver hypertrophy, kidney weight and histopathology, cellular casts in urine, thyroid hypertrophy and colloid alteration and ocular toxicity were all observed. Repeat-dose inhalation toxicity studies were not conducted and the duration of the 28-day dermal toxicity study was not appropriate for long-term exposure scenarios thus necessitating the use of an oral study for risk assessment.

The target MOE selected for this endpoint is 100. Ten-fold factors were applied each for interspecies extrapolation and intraspecies variability. This target MOE is considered to be protective of all adults including nursing infants and the unborn children of exposed female workers.

Pick-Your-Own and Residential Dermal Exposure

For the dermal risk assessment in pick-your-own and residential ornamental use scenarios, the short-term dermal toxicity study in rats was selected. At a dose of 1000 mg/kg bw/day, clinical chemistry effects and liver toxicity were evident. A NOAEL of 300 mg/kg bw/day was established.

The target MOE selected for this endpoint is 100. Ten-fold factors were applied each for interspecies extrapolation and intraspecies variability. This MOE is considered to be protective of all adults including pregnant and lactating women and their unborn children, as well as nursing infants and children of exposed women. For reasons outlined in the *Pest Control Products Act* Hazard Characterization section, the *Pest Control Products Act* factor was reduced to 1-fold.

Pick-Your-Own and Residential Oral Exposure

For the oral risk assessment in pick-your-own and residential ornamental use scenarios, the acute neurotoxicity study in rats was selected. At the doses of 100 mg/kg bw, session motor and locomotor activities were decreased in females. A NOAEL of 50 mg/kg bw was established.

The target MOE selected for this endpoint is 100. Ten-fold factors were applied each for interspecies extrapolation and intraspecies variability. This MOE is considered to be protective of all adults including pregnant and lactating women and their unborn children, as well as nursing infants and children of exposed women. For reasons outlined in the *Pest Control Products Act* Hazard Characterization section, the *Pest Control Products Act* factor was reduced to 1-fold

3.4.1.1 Dermal Absorption

An in vivo dermal absorption study in rats as well as an in vitro dermal absorption study using human and rat skin were submitted. In the in vivo rat dermal absorption study, male Wistar rats were dosed with approximately 5 or 0.005 mg/cm² fluopyram. Animals were exposed for an eight-hour period, after which time the skin was washed. Animals were terminated at 8, 24, 72 or 168 hours after dosing. The absorbed dose was calculated by summing residues in urine, faeces, cage wash, treated skin which had been tape stripped to remove the stratum corneum, surrounding skin, blood and carcass. The mean absorbable dose was 2.53, 4.53, 3.02 and 2.24% at the high dose and 12.81, 8.68, 10.96, and 11.76% at the low dose for the four termination periods, respectively.

An in vitro dermal penetration study with rat and human skin was conducted concurrently with the same doses used in the in vivo study. Human abdominal skin and rat dorsal skin were dosed in flow-through diffusion cells. Skin samples were exposed for 8 hours and then swabbed to remove non-absorbed dose. At the end of the study (24 hours), the skin was swabbed again, and then tape stripped. Radioactivity in the receptor fluid and the skin were combined to determine the absorbable dose. At the low dose, 14.76% of the applied dose was absorbable in rat skin and 2.95% of the applied dose was absorbable in human skin samples. From this study, human skin appears to be five times less permeable than rat skin.

The dermal absorption studies for fluopyram generally met the requirements and 'minimal standards' of the draft NAFTA triple pack approach (a combination of dermal absorption data including in vitro and in vivo data in rats and in vitro data in human). As such, it was considered appropriate to apply the 'triple pack' approach to this active ingredient. Due to uncertainties regarding in vitro reproducibility, variability in the in vitro human dermal absorption data and regional variability in human skin, the highest value of the human in vitro results (6.90%) was chosen instead of the mean value of the samples.

As a result, the dermal absorption value of 7% was selected for use in the risk assessment for fluopyram. This value may need to be reconsidered for formulations and uses other than those currently registered. For non-cancer risk estimates, a dermal absorption factor was not required, since the dermal toxicological endpoint was based on a dermal study.

3.4.2 Occupational Exposure and Risk

3.4.2.1 Mixer/loader/applicator Exposure and Risk Assessment

Individuals have potential for exposure to products containing fluopyram during mixing, loading and application. Exposure is expected to be of short- to intermediate-term in duration and to occur by the dermal and inhalation routes. Application is by groundboom field sprayer, airblast applicator, drip irrigation and aerial application.

Chemical-specific data for assessing human exposures during pesticide handling activities were not submitted. Exposure estimates for mixers, loaders, applicators (M/L/A) are based on data from the Pesticide Handlers Exposure Database (PHED). PHED version 1.1 is a compilation of generic mixer/loader and applicator passive dosimetry data with associated software which facilitates the generation of scenario-specific exposure estimates. With a few exceptions, the PHED estimates meet criteria for data quality, specificity and quantity outlined under the North American Free Trade Agreement Technical Working Group on Pesticides. To estimate exposure for each use scenario, appropriate subsets of A and B were created from the liquid mixer/loader and groundboom, airblast or aerial applicator database files of PHED. All data were normalized for kg of active ingredient handled. Exposure estimates are presented on the basis of the best-fit measure of central tendency (summing the measure of central tendency for each body part which is most appropriate to the distribution of data for that body part). Inhalation exposures were based on light inhalation rates (17 LPM). The exposure estimates are based on M/L/A wearing long-sleeved shirts, long pants and chemical resistant gloves (Appendix I, Table 7).

For non-cancer exposure, the maximum application rate was combined with the unit exposures and default area treated per day values. Exposure was calculated using the following equation:

Exposure = Unit Exposure (μg/kg a.i. handled) × Application Rate (kg a.i./ha) × Area Treated (ha)
(μg/kg bw/day)

Body Weight (kg)

Risk of concern is based on the equation, NOAEL/exposure, where concerns are identified if the MOE is less than the target MOE. Dermal MOEs were calculated based on a NOAEL of 300 mg/kg bw/day from a 28-day rat dermal toxicity study. Inhalation MOEs were based on a NOAEL of 12.5 mg/kg bw/day from a 90-day rat oral toxicity study. The target MOE for both routes of exposure is 100. Non-cancer exposure and risk estimates for fluopyram are presented in Appendix I, Table 8. Non-cancer MOEs for all scenarios are above the target MOE.

A deterministic cancer risk assessment was conducted for farmers and custom applicators mixing/loading and applying products containing fluopyram to the approved crops. Absorbed average daily doses (ADD; equivalent to the exposure estimate for the calculations of non-cancer MOEs with a 7% dermal absorption factor) were used as the basis for calculating lifetime average daily dose (LADD) values. Dermal and inhalation ADD values were added to obtain combined ADD values. LADD values were then calculated by amortizing exposure over the lifetime of the worker based on the use pattern using the following equation.

 $LADD = \underbrace{ADD \times Treatment\ Frequency \times Duration\ of\ Exposure\ (40\ years)}_{365\ days/year \times Life\ Expectancy\ (75\ years)}$

The treatment frequency for farmers was assumed to be equal to the maximum number of applications per year for farmers and up to 60 days per year for custom applicators, since custom applicators can apply the same product on several farms. An exposure-duration of 40-years was assumed for farmers and custom applicators.

Cancer risks were calculated by multiplying an estimated LADD by a Q_1^* for fluopyram derived from the dose response data in the appropriate toxicological study [$Q_1^* = 1.72 \times 10^{-2}$ (mg/kg bw/day)⁻¹].

Cancer Risk = LADD \times Q₁*

Cancer risks for farmers and custom applicators mixing/loading and applying products containing fluopyram to all approved crops are below 1×10^{-5} (Appendix I, Table 9), and are considered acceptable.

3.4.2.2 Exposure and Risk Assessment for Workers Entering Treated Areas

There is potential for workers entering treated fields to perform routine re-entry activities to be exposed to residues of fluopyram on foliage. Exposure is expected to be of short- to intermediate-term in duration and to occur primarily by the dermal route.

Since no chemical specific dislodgeable foliar residue (DFR) data was submitted, a default DFR value of 20% of the application rate with a 10% daily dissipation rate was used to estimate risk to workers contacting treated foliage. A tier one approach was used, in that, the highest transfer coefficient for each crop group was used to estimate exposure. Postapplication exposure was calculated using the following equation:

Exposure = <u>DFR × Transfer Coefficient × Exposure Duration (8 hours)</u> Body Weight (kg)

Non-cancer risks for workers entering treated fields for fluopyram are above the target MOE for all crops and activities (Appendix I, Table 10).

A deterministic cancer risk assessment was conducted for fluopyram for workers entering fields treated with fluopyram to all approved crops. ADD was used as the basis for calculating LADD values. A time weighted average DFR value over a 30-day period assuming 2 applications made 7 days apart, and assuming a dissipation rate of 10% per day was used in the calculation of ADD for workers entering treated areas. LADD values were then calculated by amortizing exposure over the lifetime of the worker based on the use pattern using the following equation.

LADD = ADD × Exposure Frequency × Duration of Exposure (40 years) 365 days/year × Life Expectancy (75 years)

The exposure frequency was assumed to be equivalent to 30 days for all approved crops. An exposure-duration of 40 years was assumed for re-entry workers.

Cancer risks were calculated by multiplying an estimated Lifetime LADD by a Q_1^* for fluopyram derived from the dose response data in the appropriate toxicological study $[Q_1^* = 1.72 \times 10^{-2} \text{ (mg/kg bw/day)}^{-1}]$.

Cancer Risk = LADD \times Q₁*

Cancer risks for worker entering fields treated with fluopyram are below 1×10^{-5} (Appendix I, Table 11) and are considered acceptable for all crops except wine grapes. For workers hand harvesting, training, thinning, hand pruning, tying and leaf pulling in grapes, a cancer risk estimate of 1.6×10^{-5} was calculated. This value was calculated with a 30-day time weighed average DFR value for grapes assuming two applications made 7 days apart (which is assumed to be the minimum re-treatment interval for grapes). This cancer risk estimate assumed postapplication exposure would occur daily for 8 hours per day, for 30 consecutive days following the first application, each year for 40 years. In addition, default DFR values (20% of the application rate) and a 10% daily dissipation rate were used to estimate cancer risk, and a preharvest interval (PHI) of 7 days is required for harvesting grapes. For these reasons, the cancer risk for grapes is expected to be a conservative estimate and is considered acceptable.

3.4.3 Residential Exposure and Risk Assessment

3.4.3.1 Handler Exposure and Risk

There are no domestic products; therefore no residential mixer/loader/applicator risk assessment is required.

3.4.3.2 Postapplication Exposure and Risk

There is potential for postapplication exposure to the general population entering areas treated with fluopyram. Since fluopyram is for use on apples and strawberries, exposure from pick-your-own (PYO) farms was considered as well as exposure to apple trees in residential areas. The postapplication risk assessment for workers is considered adequate to cover off risk to the general population picking apples and strawberries at PYO facilities and those exposed to treated residential apple trees since the duration of exposure is expected to be shorter than for commercial workers.

As there is potential for a person to be exposed through contact with treated foliage as well as eating the fruits that they are harvesting, both dermal and dietary exposure are generally aggregated in a PYO risk assessment. However, since no specific overlapping effects were noted between the dermal and oral endpoints chosen for fluopyram, an aggregate assessment for PYO scenarios was not required.

3.4.3.3 Bystander Exposure and Risk

Bystander exposure should be negligible since the potential for drift is expected to be minimal. Application is limited to agricultural crops only when there is low risk of drift to areas of human habitation or activity such as houses, cottages, schools and recreational areas, taking into consideration wind speed, wind direction, temperature inversions, application equipment and sprayer settings.

3.5 Food Residues Exposure Assessment

3.5.1 Residues in Plant and Animal Foodstuffs

The residue definition for enforcement is fluopyram in plant commodities, and fluopyram including the metabolite fluopyram-benzamide (expressed as parent equivalent) in animal commodities. The residue definition for risk assessment is fluopyram including the metabolite fluopyram-benzamide in crops of Crop Group 6 (Legume Vegetables) and 20 (Oilseeds), and fluopyram in all other plant commodities. The residue definition for risk assessment is fluopyram including the metabolites fluopyram-benzamide and fluopyram-olefines (total of 2 isomers) (expressed as parent equivalent) in poultry tissues and eggs, and fluopyram including the metabolites fluopyram-benzamide, fluopyram-olefines (total of 2 isomers) and fluopyram-7-hydroxy (expressed as parent equivalent) in ruminant tissues and milk.

The HPLC-MS/MS enforcement analytical methods are valid for the quantitation of fluopyram residues in crop matrices, and for the quantitation of fluopyram and the benzamide metabolite in livestock matrices. The residues of fluopyram and the benzamide metabolite are stable in representative matrices from five different crop categories (commodities with high water, high oil, high protein, high starch and high acid content) for up to 36 months when stored at -20°C. Therefore, fluopyram residues are considered stable in all frozen crop matrices and processed crop fractions for up to 36 months. Fluopyram residues concentrated in the following processed commodities: sugar beet refined sugar (1.3×), wheat bran (2.7×), wheat germ (2.4×), corn bran (2.6×), refined corn oil (2.6×) and refined peanut oil (1.5×). Adequate feeding studies were carried out to assess the anticipated residues in livestock matrices resulting from the currently approved uses. Supervised residue trials conducted throughout the United States and Canada using end-use products containing fluopyram in or on potatoes, sugar beets, dry beans, dry peas, soybeans, watermelon, apples, cherries, strawberries, wine grapes, almonds, pecans, wheat, sorghum, corn (field and sweet), canola, peanuts and cottonseed, and in Latin America on bananas are sufficient to support the proposed maximum residue limits (MRLs).

3.5.2 Exposure from Drinking Water

3.5.2.1 Concentrations in Drinking Water

Estimated environmental concentrations (EECs) of fluopyram in potential drinking water sources (groundwater and surface water) were estimated using computer simulation models. An overview of how the EECs are estimated is provided in the PMRA's Science Policy Notice SPN2004-01, Estimating the Water Component of a Dietary Exposure Assessment. EECs of fluopyram in groundwater were calculated using the LEACHM model to simulate leaching through a layered soil profile over a 50-year period. The calculated concentrations using LEACHM are based on the flux, or movement, of pesticide into shallow groundwater with time. EECs of fluopyram in surface water were calculated using the PRZM/EXAMS models, which simulate pesticide runoff from a treated field into an adjacent water body and the fate of a pesticide within that water body. Pesticide concentrations in surface water were estimated in two types of vulnerable drinking water sources, a small reservoir and a prairie dugout.

A Level 1 drinking water assessment was conducted using conservative assumptions with respect to environmental fate, application rate and timing, and geographic scenario. The Level 1 EEC estimate is expected to allow for future use expansion into other crops at this application rate. Appendix I, Table 12, lists the application information and main environmental fate characteristics used in the simulations. Fifteen initial application dates for surface water, and six initial application dates for groundwater modelling between late April and late July were modelled. The models were run for 50 years for all scenarios. The largest EECs of all selected runs are reported in Appendix I, Table 13.

The EECs for chronic refined dietary exposure assessment were not acceptable. The highest EECs for Level 1 were from the dugout scenario, and hence, it was decided to model region specific crops relevant for dugout use only (Prairie region). There were two main runs for Level 2 which reflected two different intervals for application timing, and also two different crops and regions. The first was two applications of 250 g a.i./ha each at a 7-day interval (for example, watermelon) and the second was with the same applications at a 14-day interval (for example, peanut, almond). The largest Level 2 EECs for the dugout are reported in Appendix I, Table 13.

Since some toxicological data is uncertain and is undergoing further investigation, an attempt was made to estimate risks by modelling the use of fluopyram for one, two, or three years of applications. For surface water, these limited applications were tested on Saskatchewan and Prince Edward Island scenarios. For dugout modelling, the use pattern and date of application were the same as in Level 2 (2 × 250 g a.i./ha at a 14-day interval). For reservoir modelling, PEI-potato was run with the use pattern modeled at Level 1 (2 × 250 g a.i./ha at a 7-day interval). The application date selected for the runs was the date giving the highest EEC in Level 1 modelling. For groundwater, the use pattern was the same as in Level 1 (2 × 250 g a.i./ha at a 7-day interval). For each restricted use pattern, the LEACHM model was run 12 times each with fluopyram applied starting in one of the first twelve years of the simulation. This gave 12 different EEC's for each case. In addition the aerobic soil metabolism half-life was recalculated by taking the 80th percentile of a lognormal distribution fitted to the seven available values. Results for the additional Level 2 modeling are summarized in Appendix I, Table 14.

For the ground water restricted applications, further analysis was performed for consideration of chronic effects by providing EECs averaged over 5, 10, 20 and 70 year periods. These are shown in Appendix I, Table 15, together with the daily and yearly EECs. EECs for all eleven groundwater scenarios have been provided to allow for consideration of crops restrictions. Also for information purposes, the numbers of days when EECs exceed 2 μ g/L for each of the eleven scenarios are provided in Appendix I, Table 16.

Additional Level 2 modelling was conducted for groundwater. A reduced potato use rate at yearly application of 400 g a.i./ha (two applications of 150 g a.i./ha plus one of 100 g a.i./ha at the interval of 7 days) for three consecutive years application only and 100 years of consecutive application was modelled for groundwater. The groundwater EECs averaged over 70 years are reported in Appendix I, Table 17, for the three and 100 consecutive years of application.

3.5.3 Dietary Risk Assessment

Acute and chronic (cancer and non-cancer) dietary risk assessments were conducted using the Dietary Exposure Evaluation Model (DEEM–FCID™, Version 2.14), which uses updated food consumption data from the United States Department of Agriculture's Continuing Surveys of Food Intakes by Individuals, 1994–1996 and 1998.

3.5.3.1 Chronic Dietary Exposure Results and Characterization

The following criteria were applied to the refined chronic non-cancer analysis: Supervised trial mean residue (STMR) values, experimental processing factors, where available, Canadian and American projected percent crop treated values, and anticipated residues for livestock commodities. The refined chronic dietary exposure from all supported fluopyram food uses (alone) for the total population, including infants and children, and all representative population subgroups is less than 7% of the ADI. Aggregate exposure from food and water is considered acceptable. The PMRA estimates that chronic dietary exposure to fluopyram from food and water is 19.6% (0.002350 mg/kg bw/day) of the ADI for the total population. The highest exposure and risk estimate is for infants less than one year old at 63.8% (0.007661 mg/kg bw/day) of the ADI.

The refined cancer risk assessment was conducted based on a limited three-year application period and with the same criteria used for the chronic non-cancer assessment. The lifetime cancer risk from exposure to fluopyram in food and water is estimated to be 1×10^{-6} for the general population, which is considered acceptable.

3.5.3.2 Acute Dietary Exposure Results and Characterization

The following criteria were applied to the basic acute analysis: 100% crop treated, default processing factors, and residues of fluopyram in/on crop and animal commodities at MRL levels. The basic acute dietary exposure from all supported fluopyram food uses was estimated to be 4.4% of the ARfD for the general population (95th percentile, deterministic). Aggregate exposure from food and water is considered acceptable and below PMRA's level of concern. Specifically, an acute dietary exposure of 2.8% to 9.8% of the ARfD was obtained for all population subgroups, with children 1-2 years old as the highest exposed population subgroup.

3.5.4 Aggregate Exposure and Risk

The aggregate risk for fluopyram consists of exposure from food and drinking water sources only. Given that apples and strawberries can be treated with fluopyram, there is potential for exposure to fluopyram during pick-your-own harvesting activities and during harvesting of fruit from trees, in residential settings, that may have been treated. Since the acute dietary and short-term dermal toxicological endpoints are based on different toxicological effects, no aggregation of dermal and dietary exposure is required.

3.5.5 Maximum Residue Limits

Table 3.5.1 Proposed Maximum Residue Limits

Commodity	Recommended MRL (ppm)	
Wine grapes	2.0	
Canola	1.8	
Crop Group 15 (except rice) - Cereal Grains, except rice; Cherries; Strawberries	1.5	
Bananas; Watermelon	1.0	
Dry chickpeas and dry lentils	0.4	
Apples	0.3	
Sugar beet roots; Dry soybeans	0.1	
Grain lupin, dry kidney beans, dry lima beans, dry navy beans, dry pink beans, dry pinto beans, dry tepary beans, dry beans, dry adzuki beans, dry blackeyed peas, dry catjang seed, dry cowpea seed, dry moth beans, dry mung beans, dry rice beans, dry southern peas, dry urd beans, dry broad beans, dry guar seed, dry lablab beans	0.09	
Crop Group 14 – Tree Nuts Group	0.05	
Crop Subgroup 1C - Tuberous and Corm Vegetables Subgroup; Peanuts	0.02	
Undelinted cotton seeds	0.01	
Meat byproducts of cattle, goats, horses and sheep	0.40	
Meat byproducts of poultry	0.10	
Eggs; Milk	0.06	
Fat and meat of cattle, goats, horses and sheep	0.05	
Meat byproducts of hogs; Fat and meat of poultry	0.03	
Fat and meat of hogs	0.02	

MRLs are proposed for each commodity included in the listed crop groupings in accordance with the Residue Chemistry Crop Groups webpage in the Pesticides and Pest Management section of Health Canada's website.

For additional information on MRL in terms of the international situation and trade implications, refer to Appendix II.

The nature of the residues in animal and plant matrices, analytical methodologies, field trial data, and acute and chronic (cancer and non-cancer) dietary risk estimates are summarized in Appendix I, Tables 1, 18a-18h and 19.

4.0 Impact on the Environment

4.1 Fate and Behaviour in the Environment

Physico-chemical properties, fate and behaviour of fluopyram in terrestrial and aquatic systems are summarized in Appendix I, Tables 20-24.

Based on its physical and chemical properties, fluopyram is soluble in water, is not likely to volatilize from moist soil or water surfaces under field conditions, has a limited potential for phototransformation in the environment, does not dissociate under environmentally relevant pH conditions and has a potential for bioaccumulation in aquatic organisms.

Fluopyram is stable to hydrolysis, photolysis, aerobic and anaerobic biotransformation in soils. It is persistent in soils and has a potential for long-term accumulation and residue carry over to the following crop season. No major transformation products were detected in soils in laboratory and field studies under Canadian field use conditions. Minor transformation products identified in soils were fluopyram-7-hydroxy, fluopyram-pyridyl-carboxylic acid, fluopyram-benzamide and fluopyram-methyl-sulfoxide (only in laboratory studies). Fluopyram forms neither major nor minor transformation products in soils under anaerobic conditions.

Based on the laboratory adsorption studies, fluopyram is classified as moderately mobile in soils. In field studies, residues of fluopyram were detected beyond 30 cm soil depths. These studies indicate that fluopyram has a potential to leach and contaminate the groundwater depending on the soil type and location. None of the transformation products were, however, detected beyond 30 cm soil depth, which indicate that they have a low potential to leach and contaminate the groundwater. According to the bioaccumulation study with bluegill sunfish, fluopyram has a low potential for bioconcentration/bioaccumulation in organisms.

Fluopyram can enter aquatic systems through spray drift, overland runoff or through the movement of soil particles with bound residues. Photolysis is not an important route of transformation in the aquatic environment. Fluopyram is persistent in sediment/water aquatic systems under aerobic and anaerobic conditions and partitions significantly from water to the sediment. No major transformation products were detected in the water or sediment phases. Several minor transformation products were detected in natural water of which one was identified as fluopyram-lactam.

Based on relatively low vapour pressure and Henry's Law Constant, fluopyram is not expected to partition to the atmosphere.

4.2 Environmental Risk Characterization

The environmental risk assessment integrates the environmental exposure and ecotoxicology information to estimate the potential for adverse effects on non-target species. This integration is achieved by comparing exposure concentrations with concentrations at which adverse effects occur. The EECs are concentrations of pesticide in various environmental media, such as food,

water, soil and air. The EECs are estimated using standard models which take into consideration the application rate(s), chemical properties and environmental fate properties, including the dissipation of the pesticide between applications. Ecotoxicology information includes acute and chronic toxicity data for various organisms or groups of organisms from both terrestrial and aquatic habitats including invertebrates, vertebrates, and plants. Toxicity endpoints used in risk assessments may be adjusted to account for potential differences in species sensitivity as well as varying protection goals (that is, protection at the community, population, or individual level).

Initially, a screening level risk assessment is performed to identify products and/or specific uses that do not pose a risk to non-target organisms, and to identify those groups of organisms for which there may be a potential risk. The screening level risk assessment uses simple methods, conservative exposure scenarios (for example, direct application at a maximum cumulative application rate) and sensitive toxicity endpoints. Screening level EECs in soil, water, aquatic eco-scenarios, vegetation and other food sources are presented in Appendix I, Tables 25-27 and Tables 38-39.

A risk quotient (RQ) is calculated by dividing the exposure estimate with an appropriate toxicity value (RQ = exposure/toxicity), and the RQ is then compared to the level of concern (LOC). If the screening level RQ is below the level of concern (LOC = 1), the risk is considered negligible and no further risk characterization is necessary. If the screening level RQ is equal to or greater than the LOC, then a refined risk assessment is performed to further characterize the risk. A refined assessment takes into consideration more realistic exposure scenarios (such as drift to non-target habitats) and might consider different toxicity endpoints. Refinements may include further characterization of risk based on exposure modelling, monitoring data, results from field or mesocosm studies, and probabilistic risk assessment methods. Refinements to the risk assessment may continue until the risk is adequately characterized or no further refinements are possible.

4.2.1 Risks to Terrestrial Organisms

A risk assessment of fluopyram and its associated end-use products was undertaken for terrestrial organisms based on available toxicity data for earthworms (acute and chronic), bees (acute oral and contact), predatory and/or parasitic invertebrates, birds (acute oral, dietary and chronic), mammals (acute oral, dietary and chronic) and terrestrial plants (effects on seedling emergence and vegetative vigour). A summary of terrestrial toxicity data for fluopyram is presented in Appendix I, Table 28, and the accompanying screening level risk assessment in Appendix I, Tables 29, 30, 31, 33, 34, 36, 40, 41, 43, 44, 46, 47, 48 and 50. Refined EECs and risk assessments for fluopyram with spray drift and runoff water are presented in Appendix I, Tables 32, 35, 37, 42, 45 and 49.

Earthworm

Fluopyram is not acutely toxic to earthworms. Although chronic effects (reproduction) were observed, low RQ values indicated that the LOC was not exceeded and, therefore, fluopyram will not pose a risk to earthworms.

Honey bees

No mortalities or adverse effects were observed when bees were exposed to fluopyram on an acute oral or contact basis. The low RQ values indicated that the level of concern was not exceeded and, therefore, fluopyram will not pose a risk to bees.

Parasitic wasps and predatory mites

No acute toxicity was observed in wasps and mites when exposed to fluopyram. Although chronic effects (reproduction) were observed in mites, the low RQ values indicated that the LOC was not exceeded and therefore, fluopyram will not pose a risk to parasitic wasps and predatory mites.

Wild birds and mammals

To characterize exposure, the concentration of fluopyram on various food items is used to determine the amount of pesticide in the diet, or estimated daily exposure (EDE). As exposure is dependent on the body weight of the organism and the amount and type of food consumed, a set of generic body weights is used to represent a range of species (20, 100 and 1000 g for birds, and 15, 35 and 1000 g for mammals). In addition, specialized feeding guilds are considered for each category of animal weights (herbivore, frugivore, insectivore and granivore). The EDE is calculated as follows: EDE = (FIR/bw) × EEC, where the food ingestion rate (FIR) is based on equations from Nagy (1987), bw is the generic body weight of the organism, and the EEC is the expected environmental concentration.

At the screening level, the risk is characterized only for feeding guilds associated with the most conservative exposure estimate (insectivores feeding on small insects or herbivores feeding on short grass) and it is assumed that food items are contaminated with maximum residue levels. In addition, only acute oral and reproduction endpoints are considered.

Wild birds: Fluopyram is not acutely toxic to birds. The RQ values were less than the LOC and small, medium and large birds are, therefore, not at potential risk on an acute basis.

Fluopyram adversely affects the reproductive performance of birds if the level of consumption exceeds 4.12 mg a.i./kg bw/day. Screening level risk assessment indicated that fluopyram may pose a risk to reproductive performance of small, medium and large birds.

The risk assessment for reproduction was therefore expanded to include all relevant food guilds and food items and also to include both on-field and off-field exposure scenarios with both maximum and mean nomogram residue concentrations. For off-field scenarios, a percent drift of 74, 59 and 6% was considered for early airblast, late airblast and field spray applications, respectively.

When considering even mean nomogram residues, on-field RQs exceeded the LOC for small and medium insectivores for all three end-use products, as well as for large herbivores with Luna Privilege. Off-field RQs exceeded the LOC only for airblast applications with Luna Privilege and Luna Tranquility Fungicide. The highest RQs were observed for small insectivorous birds for both on-field and off-field scenarios.

To further explore the potential for reproductive concern, a refined risk assessment was undertaken based on the LOAEL. The RQ values still exceeded the LOC for small and medium insectivorous birds with exposure to mean concentrations; these RQs, however, only marginally exceeded the LOC. Due to the conservative nature of the risk assessment, these marginal exceedances of the LOC are unlikely to result in adverse effects on reproductive performance. However, as a precautionary measure, bird toxicity label statements are required.

Wild mammals: Fluopyram is acutely non-toxic to mammals. The RQ values were less than the LOC and the small, medium and large mammals are, therefore, not at potential risk on an acute basis.

Fluopyram adversely affects the reproductive performance of mammals if the level of exposure exceeds 13.9 mg a.i./kg bw/day. The reproductive RQ values with direct exposure to contaminated food in the treated field (on-field) exceeded the LOC for medium and large sized mammals with Luna Privilege and medium sized mammals with Luna Tranquility Fungicide. The risk assessment for reproduction was, therefore, expanded to include all relevant food guilds and food items including on-field and off-field exposure scenarios with maximum and mean residue concentrations. For off-field scenarios, a percent drift of 74, 59 and 6% was considered for early airblast, late airblast and field spray applications, respectively.

Risk quotients exceeded the level of concern only for medium and large herbivores when considering exposure with maximum residue concentrations. With mean residue concentrations, the LOC was, however, not exceeded for any of the feeding guilds. Also, no reproductive risk was identified for all mammals feeding on food items contaminated from spray drift off the treated area (field spray applications).

The risk assessment was based on the conservative assumption that a mammal fed on 100% of a given food item and that all food was contaminated. Given that the LOCs were exceeded by a small margin for some but not for all food items considered in the risk assessment, the overall risk to mammals is considered to be low and that the wild mammals are likely have a diet comprised of different types of food items. To further support this conclusion, reproduction RQs were also calculated using a LOAEL. The RQ values with the LOAEL indicate that the LOC was not exceeded for both on-field and off-field exposure to maximum as well as mean residue concentrations for medium sized mammals.

As such, the risk to reproductive performance of wild mammals is expected to be limited (minimal).

Non-target terrestrial plants

Studies on toxicity/effects on seedling emergence and vegetative vigour indicated EC₂₅ values of greater than 500 and 250 g a.i./ha (the highest applications rates tested), respectively. The RQ values exceeded the LOC with Luna Privilege, but not with the Luna Tranquility Fungicide and Propulse Fungicide. As such, fluopyram may affect plant growth with the approved uses of Luna Privilege.

As screening level risk assessment indicated a risk, a refined risk assessment was undertaken to assess the risk to non-target plants due to spray drift. Three application scenarios (airblast early (74% drift), airblast late (59% drift) and ground boom (6% drift) applications) were used to assess the risk to non-target plants due to spray drift.

As the RQ values indicated that the LOC was not exceeded, the approved uses of Luna Privilege will not affect the seedling emergence with all the three application scenarios. For vegetative vigour, however, the RQ value is slightly greater than one for the airblast early application scenario and, therefore, the approved uses of Luna Privilege may pose a risk to non-target terrestrial plants. Risk mitigation measures such as buffer zones are, therefore, required to protect non-target terrestrial habitats.

4.2.2 Risks to Aquatic Organisms

Aquatic organisms can be exposed to fluopyram as a result of spray drift and over-land run-off. To assess the potential for adverse effects, screening level EECs in the aquatic environment based on a direct application to water were used as exposure estimates. A risk assessment of fluopyram end-use products was undertaken for freshwater and marine aquatic organisms based on available toxicity data for algae (acute), aquatic plants (acute), invertebrates (acute and chronic), fish (acute and chronic) and amphibians (using fish as surrogate data).

A summary of aquatic toxicity data for fluopyram is presented in Appendix I, Table 51. For acute toxicity studies, uncertainty factors of 1/2 and 1/10 EC(LC)₅₀ are used in modifying the toxicity values for aquatic plants and invertebrates, and fish species, respectively when calculating RQs. No uncertainty factors are applied to chronic NOEC endpoints. For groups where the LOC is exceeded (that is, RQ \geq 1), a refined Tier 1 assessment is conducted to determine risk resulting from spray drift and runoff water separately. The calculated RQs are summarized in Appendix I, Tables 52 and 56 (screening level), 53 & 57 (Tier 1 runoff) and 54 and 58 (Tier 1 spray drift).

Freshwater fish

Fluopyram is acutely toxic to cold and warm water fish and also would result in chronic adverse effects at concentrations greater than 0.135 mg a.i./L. The low RQ values, however, indicate that the LOC was not exceeded and therefore, the freshwater fish are not a potential risk. Further, a bioaccumulation study with bluegill sunfish showed that fluopyram has a low potential for bioconcentration/bioaccumulation in aquatic organisms.

Amphibians

As no amphibian data were submitted, acute and chronic risk to amphibians were assessed using surrogate values of the most sensitive fish species – that is, rainbow trout and fathead minnow, respectively. The EECs for ground application were estimated for a water depth of 15 cm. The RQ values for the acute and chronic exposures exceeded the LOC, which indicate that the approved uses of fluopyram may pose acute and chronic risks to amphibians.

The Screening level risk assessment conducted was a conservative scenario of direct application into a body of water. As this assessment indicated a potential risk to amphibians, a refined risk assessment was conducted by estimating EECs in runoff water from treated areas into a receiving water body and by spray drift.

For acute risk to amphibians from runoff, the estimated peak EEC (acute exposure in a 15 cm depth water body) from the aquatic eco-scenario modelling was used to assess the acute risk. The acute LC₅₀ for the most sensitive fish species, rainbow trout, was used as a surrogate for the amphibians. The RQ values indicated that the LOC was still exceeded and therefore, the approved of uses of fluopyram may pose an acute risk for amphibians from runoff.

For chronic risk to amphibians from runoff, the estimated EEC (21 day chronic exposure in a 15 cm depth water body) from the aquatic eco-scenario modelling was used for the risk assessment. The 21-day EEC was chosen to calculate the RQ as the chronic fathead minnow study period was 33 days. The chronic NOEC for fathead minnow was used as a surrogate for the amphibians. The RQ value indicated that the LOC was exceeded and the approved uses of fluopyram may pose a chronic risk for amphibians.

Three application scenarios, airblast early (74% drift), airblast late (59% drift), and ground boom (6% drift) applications were used to assess the risk to amphibians due to spraydrift. The acute and chronic RQs values indicated that the LOC was exceeded and the approved uses of fluopyram may pose an acute and chronic risk for amphibians due to spray drift from airblast early and late applications. The LOC was not exceeded for ground boom applications.

A refined risk assessment with run-off and spray drift (airblast) scenarios indicated that the approved uses of fluopyram may pose a risk to amphibians and, therefore, risk mitigation measures such as buffer zones are required to protect these organisms.

A screening level risk assessment was conducted with an EEC from direct aerial overspray with the approved application rates for potatoes. This assessment indicated that the LOC was exceeded for acute and chronic exposures and, therefore, a refined risk assessment was conducted with 23% spray drift for aerial applications (Appendix I, Table 55). This assessment indicated that the LOC was not exceeded for acute and chronic exposures for one metre off-field and, therefore, amphibians in the off-field are not at risk from the approved aerial applications for potato. A default buffer zone of one meter is, however, approved to cover uncertainty between direct overspray and one meter off field exposure.

Freshwater invertebrates

Fluopyram is acutely toxic to freshwater invertebrates (*Daphnia* sp.) and would result in adverse chronic effects if the concentrations in water exceed 1.214 mg a.i/L. The acute and chronic RQ values, however, indicated that the LOC was not exceeded and, therefore, the approved uses of fluopyram would pose a negligible risk to freshwater aquatic invertebrates.

Sediment-dwelling organisms

Fluopyram is persistent in aquatic systems and therefore, risk to sediment-dwelling organisms was also assessed. Chronic toxicity data for *C. riparius* and *C. tentans* were submitted which indicate that chronic adverse effects would result if the concentrations in sediment and pore water exceed 26.0 and 3.8 mg a.i./L, respectively. The RQ values were less than the LOC which indicated that the approved uses of fluopyram would pose a negligible risk to sediment-dwelling organisms.

Freshwater algae

Fluopyram is acutely toxic to freshwater algae and the most sensitive freshwater algal species is green algae. Low RQ values, however, indicated that the LOC was not exceeded and, therefore, the approved uses of fluopyram would pose a negligible risk to freshwater algae.

Freshwater plants

Adverse effects on aquatic plant, *Lemna gibba*, were observed when exposed to fluopyram. Low RQ values, however, indicated that the LOC was not exceeded and, therefore, the approved uses of fluopyram would pose a negligible risk to aquatic plants.

Marine fish

Fluopyram is acutely toxic to marine fish and the most sensitive species is sheepshead minnow. The RQ value, however, indicated that the LOC was not exceeded and therefore, marine fish are not at potential risk with the approved uses of fluopyram.

Marine invertebrates

Fluopyram is acutely toxic to marine invertebrates and the most sensitive species is eastern oyster. The low RQ values, however, indicated that the LOC was not exceeded and therefore, marine invertebrates are not at potential risk with the approved uses of fluopyram.

Marine algae

Fluopyram is acutely toxic to marine algae and the most sensitive species is saltwater diatom. The low RQ values, however, indicate that LOC was not exceeded and therefore, marine algae are not at potential risk with the approved uses of fluopyram.

Marine amphipods

Fluopyram is acutely toxic to marine amphipods and would result in chronic adverse effects if exposed to concentrations greater than 0.55 mg a.i./L. The low acute and chronic RQ values, however, indicated that the LOC was not exceeded and therefore, the approved uses of fluopyram would pose a negligible risk to marine amphipods.

5.0 Value

5.1 Effectiveness Against Pests

5.1.1 Acceptable Efficacy Claims

The number of submitted trials reviewed in support of the efficacy claims on the Luna Privilege, Luna Tranquility Fungicide, and Propulse Fungicide labels totalled 57, 18, and 14 trials, respectively. Proposed and supported claims are listed in Appendix I, Tables 59 to 61.

5.1.1.1 Almond and cherry

Brown rot blossom blight

The same five trials were used as evidence to support the efficacy of Luna Privilege against brown rot blossom blight in both almond and cherry given the similarities between the two crops and their susceptibility to the disease. Four trials were conducted on various related species of stone fruit trees including cherry. The fifth trial was conducted on almond. Averaged across the different trials, disease severity reduction reached up to 84%. Average levels of reductions in disease incidence across the trials were somewhat lower at 61%. Although almond production in Canada is negligible, developments in almond breeding have opened the possibility of introductions of hardier types and varieties of almonds that could be used in establishing viable commercial production in Canada.

5.1.1.2 Apple

Leaf scab

Five trials conducted on apple were used to demonstrate the efficacy of Luna Privilege against leaf scab. Up to 100% control, in terms of both disease severity and incidence, was obtained on leaves of tested trees. Disease control on fruits, although more variable than on leaves, also reached levels up to 100% in certain trials. It was noted that the label claim is indicated for control of the disease specifically on the leaf, rather than the fruit. Therefore, the levels of disease control observed across the apple trials were sufficient to support the claim for control of leaf scab.

In addition, Luna Tranquility Fungicide was also shown to be highly effective against this disease. Across the nine trials where the co-formulation was tested on apple leaf scab, up to 83% control was observed when fluopyram and pyrimethanil were applied together. Applied individually, each active ingredient provided significantly lower levels of protection than the combination product. This, along with considerations related to disease resistance management, further demonstrated the value of the co-formulation.

Powdery mildew

In two trials, Luna Tranquility Fungicide provided high levels of protection and performed better than a registered standard. Control, in terms of disease severity, was reported to have reached almost 100% in the Luna Tranquility Fungicide treatment. Although pyrimethanil was shown to have limited activity against powdery mildew in apple on its own, the main benefits of the coformulation are primarily in broadening the spectrum of diseases controlled.

5.1.1.3 Bean, dry

Powdery mildew

Direct evidence used in demonstrating Luna Privilege efficacy against powdery mildew in legume vegetables, including dry bean, was obtained from one trial conducted on peas. This evidence was supplemented by trials conducted on other crops (for example, cucurbits and wheat) where powdery mildew is caused by different but related species of pathogens. In all of these trials, Luna Privilege provided excellent protection against powdery mildew. Specifically, in the pea trial, disease severity was reduced by 81 to 100% under high disease pressure.

White mold

Across three trials on dry bean and one trial on edible bean, which was accepted as support for the dry bean claim, severity and incidence of white mold were generally reduced by over 90% in stems and pods by Luna Privilege treatments. These levels of efficacy were sufficient in demonstrating acceptable levels of white mold control in dry bean.

The combined efficacy of the two active ingredients in Propulse Fungicide was tested in six trials. Damage caused by white mold along with infection of pods and yields were assessed in the various trials. In one of the trials, reductions in the percentage of damage caused by white mold reached up to 98% under moderate to high disease pressure. Higher yields relative to the untreated control were observed in the plots treated with the co-formulation in all of the trials where yield was measured. In another trial, where the percentage of infected pods was assessed, reductions of up to 70% by Propulse Fungicide relative to the untreated control plots were observed. Overall, the product provided equivalent or superior protection to the tested standard currently registered for control of white mold in dry bean.

Ascochyta blight & mycosphaerella blight

Efficacy of Luna Privilege against ascochyta blight on dry bean was demonstrated across six field trials conducted on chickpea and lentil. In most trials, Luna Privilege was shown to be considerably more effective at reducing levels of disease severity than disease incidence. Under very high disease pressure, disease control reached up to 86% in one of the chickpea trials.

Because mycosphaerella blight is closely related to ascochyta blight and both diseases are biologically similar, the evidence described above was also deemed to be supportive of the mycosphaerella blight claim. In addition, two trials on pea directly assessing the effect of Luna Privilege on mycosphaerella blight demonstrated similarly high levels of protection under high disease pressure.

The combined effect of the two active ingredients in the co-formulated product Propulse Fungicide on ascochyta blight and mycosphaerella blight was demonstrated across eight trials on lentil, pea, and chickpea. As with fluopyram alone, the combination of two active ingredients provided excellent levels of reduction in disease severity. For instance, over 81% control of ascochyta blight was obtained in the lentil trial where the higher of the two label rates of Propulse Fungicide was applied. In five different trials conducted on chickpea, damage caused by ascochyta blight was reduced by an average of 85% and 91% when assessed 12 to 30 days after the final application of Propulse Fungicide at the low and high label rates, respectively. It was also observed that, on average, both label rates of Propulse Fungicide provided substantially higher levels of protection than the tested commercial standards in terms of disease severity and damage.

As both active ingredients in Propulse Fungicide are known to be effective on their own, either from previously registered claims or from the fluopyram trials described above, it can be concluded that this product provides added benefits in terms of resistance management.

5.1.1.4 Cherry

Powdery mildew

Two trials were conducted to demonstrate the efficacy of Luna Privilege against powdery mildew on cherry. Under heavy infestation, applications of the product at the lower labelled rate provided 62% reductions in disease. However, under moderate disease pressure and at the higher labelled rate, disease control reached 94% relative to the untreated control treatment. As described for other crops appearing on the label, Luna Privilege also has demonstrated excellent efficacy against a number of other powdery mildew-causing organisms.

5.1.1.5 Grape, wine

Powdery mildew

The efficacy of the combination product Luna Tranquility Fungicide was tested in four trials. Under moderate to high disease pressure, the product provided consistently high levels of control. Reductions in disease severity and incidence both reached 100% in many instances. Although the efficacy of fluopyram alone was not tested directly in these trials, it was indirectly demonstrated by observations where the combination of fluopyram and pyrimethanil provided 100% control where pyrimethanil alone provided a maximum of 36% control under high disease pressure. In light of demonstrated efficacy for both components, the combination product offers the benefit of simultaneous applications of multiple modes of fungicide action that are effective, thereby reducing the risk of resistance development.

Botrytis bunch rot/Grey mold

Across four trials conducted on grape, Luna Privilege provided excellent levels of protection against botrytis bunch rot. Control, in terms of disease severity, ranged from 83-99% and was shown to be equivalent to currently registered standards under moderate and high disease pressure. Reductions of disease incidence were also high, ranging from 58-92%. As efficacy of fluopyram applied alone was demonstrated in these trials and pyrimethanil, as the lone active ingredient of Scala SC Fungicide, is already registered for the control of botrytis bunch rot, the combination product Luna Tranquility Fungicide is expected to provide dual effective modes of action against botrytis bunch rot and reduce the risk of resistance development.

5.1.1.6 Peanut

Early and late leaf spot

Luna Privilege showed excellent levels of early leaf spot control in three field trials conducted on peanut. Control, in terms of disease incidence and severity, reached 89% and 96%, respectively. On the other hand, the product's efficacy against late leaf spot was demonstrated in two other field trials. In these, incidence and severity of late leaf spot were both reduced by more than 80%.

5.1.1.7 Potato

Early blight

Luna Privilege provided excellent levels of early blight control across four trials conducted on potato. Under moderate to high disease pressure, early blight severity and incidence was reduced by up to 90% and 100%, respectively. In addition, aerial applications were shown to be equally as effective as ground applications of Luna Privilege.

5.1.1.8 Strawberry

Powdery mildew

A total of six trials were conducted to demonstrate the efficacy of Luna Privilege in reducing powdery mildew in strawberry. Only applications by chemigation appear on the label. Luna Privilege provided high levels of efficacy against powdery mildew when applied to strawberry. The labelled rate of Luna Privilege provided average reductions of disease severity and incidence across the three chemigation trials of around 72% and 70%, respectively. Maximum levels of disease reduction reached 82% and 93% for severity and incidence of powdery mildew, respectively.

5.1.1.9 Watermelon

Powdery mildew

Thirteen trials demonstrating efficacy on two different species of powdery mildew-causing organisms conducted on a variety of cucurbit crops (pumpkin, squash, melon, cucumber, and zucchini) were reviewed as evidence for this claim. This set of data provided excellent support for Luna Privilege efficacy against both species of powdery mildew. The highest reductions in disease severity across the trials ranged from 81-100%, all under at least moderate, and often high disease pressure.

Botrytis grey mold

In trials conducted on other crops (grape and strawberry), high levels of protection by Luna Privilege were demonstrated against the same pathogen that causes grey mold in cucurbits. Because of the similarities in grey mold susceptibility shared among the tested crops and cucurbits, the results of these trials were extrapolated as evidence to support this claim on watermelon.

5.2 Phytotoxicity

Observations of crop tolerance to Luna Privilege, Luna Tranquility Fungicide, and Propulse Fungicide, were reported in a total of 61, 23, and 16 submitted field trials, respectively. Phytotoxicity was not observed from any of the three products when applied at rates consistent with their labelled use patterns.

5.3 Economics

No market analysis was done for this submission.

5.4 Sustainability

5.4.1 Survey of Alternatives

The chemical and other non-conventional/biological fungicidal active ingredients listed in Appendix I, Table 62, are found in products that are registered for control or suppression of diseases indicated on the Luna Privilege, Luna Tranquility Fungicide, and Propulse Fungicide labels.

5.4.2 Compatibility with Current Management Practices Including Integrated Pest Management

The use of fluopyram products should be integrated into a disease management program to attenuate the probability of resistance development to fungicides that have a similar mode of action. Integrated pest management (IPM) promotes the integration of cultural, biological, mechanical and chemical control strategies. Proper use of IPM aims to reduce pesticide use while maintaining economic returns through effective pest control and maximum crop production. Fluopyram fungicides represent one component of the chemical strategy for disease control. The use of fluopyram will complement other disease management strategies in the supported crops.

The addition of fluopyram as another chemical control option will potentially increase the longevity of other products with different modes of action as viable options for specific disease control. Combining chemical control with other cultural or biological control measures should minimize the dependence on any one control measure and therefore minimize the potential for resistance or increased tolerance to develop to any one control measure.

5.4.3 Information on the Occurrence or Possible Occurrence of the Development of Resistance

Fungicides in the succinate dehydrogenase inhibitors group, such as fluopyram, are considered to present a medium to high risk of disease resistance development by Fungicide Resistance Action Committee. Resistance to this group of fungicides has been observed for several fungal species in field populations and lab mutants. Among cases of field resistance reported by Fungicide Resistance Action Committee that are specifically relevant to the uses registered on the three fluopyram-containing product labels are resistant field isolates of powdery mildew in cucurbits and botrytis in various hosts. In addition, suspected resistant isolates of *Sclerotinia sclerotiorum*, the pathogen that causes white mold on bean and other legumes, were found in European rape seed fields.

5.4.4 Contribution to Risk Reduction and Sustainability

Luna Privilege, Luna Tranquility Fungicide, and Propulse Fungicide are safe on labelled crops and fit well into current IPM strategies when used according to directions. These broad spectrum products will benefit fruit and vegetable producers and offer a useful alternative in disease resistance management. In addition, because Luna Tranquility Fungicide and Propulse Fungicide each combine two active ingredients with different modes of action, the risk of disease resistance development is reduced in targeted pathogens that are sensitive to the two active ingredients.

6.0 Pest Control Product Policy Considerations

6.1 Toxic Substances Management Policy Considerations

The Toxic Substances Management Policy (TSMP) is a federal government policy developed to provide direction on the management of substances of concern that are released into the environment. The TSMP calls for the virtual elimination of Track 1 substances [those that meet all four criteria outlined in the policy: in other words, persistent (in air, soil, water and/or sediment), bio-accumulative, primarily a result of human activity and toxic as defined by the *Canadian Environmental Protection Act*].

During the review process, fluopyram and its transformation products were assessed in accordance with the PMRA Regulatory Directive DIR99-03⁴ and evaluated against the Track 1 criteria. The PMRA has reached the following conclusions:

- Fluopyram does not meet all Track 1 criteria, and is not considered a Track 1 substance.
 (Appendix I, Table 63)
- Fluopyram does not form any transformation products that meet all Track 1 criteria.

Technical grade fluopyram and its associated end-use products do not contain any formulants or contaminants of health or environmental concern identified in the Canada Gazette.

The use of formulants in registered pest control products is assessed on an ongoing basis through PMRA formulant initiatives and Regulatory Directive DIR2006-02.⁵

DIR99-03, The Pest Management Regulatory Agency's Strategy for Implementing the Toxic Substances
Management Policy

⁵ DIR2006-02, Formulants Policy and Implementation Guidance Document.

6.2 Formulants and Contaminants of Health or Environmental Concern

During the review process, contaminants in the technical and formulants and contaminants in the end-use products are compared against the *List of Pest control Product Formulants and Contaminants of Health or Environmental Concern* maintained in the *Canada Gazette*.⁶. The list is used as described in the PMRA Notice of Intent NOI2005-01⁷ and is based on existing policies and regulations including DIR99-03 and DIR2006-02, and taking into consideration the Ozone-depleting Substance Regulations, 1998, of the *Canadian Environmental Protection Act* (substances designated under the Montreal Protocol). The PMRA has reached the following conclusions:

The end-use products have, as a component, the preservative 1,2-benzisothiazoline-3-one (0.015%), which contains low levels of polychlorinated dibenzodioxins and furans (TSMP Track 1). As the use of this preservative was recently re-evaluated and found to be acceptable, and because the input of dioxins into the environment from pesticides is being managed as outlined in the PMRA Regulatory Directive DIR99-03 for the implementation of TSMP, the Agency position is that no further action is required.

The use of formulants in registered pest control products is assessed on an ongoing basis through PMRA formulant initiatives and Regulatory Directive DIR2006-02.

7.0 Summary

7.1 Human Health and Safety

The toxicology database submitted for fluopyram is adequate to define the majority of toxic effects that may result from exposure. There was no evidence of increased susceptibility of the young in reproduction or developmental toxicity studies. While motor/locomotor activity were decreased in the neurotoxicity study, fluopyram is not believed to be selectively neurotoxic. In short-term and chronic studies on laboratory animals, the primary targets were the liver, thyroid and kidneys. Although fluopyram was not genotoxic, there was evidence of oncogenicity in mice and rats after chronic dosing. The risk assessment protects against the toxic effects noted above by ensuring that the level of human exposure is well below the lowest dose at which these effects occurred in animal tests.

Canada Gazette, Part II, Volume 139, Number 24, SI/2005-114 (2005-11-30) pages 2641–2643: List of Pest Control Product Formulants and Contaminants of Health or Environmental Concern and in the order amending this list in the Canada Gazette, Part II, Volume 142, Number 13, SI/2008-67 (2008-06-25) pages 1611-1613. Part 1 Formulants of Health or Environmental Concern, Part 2 Formulants of Health or Environmental Concern that are Allergens Known to Cause Anaphylactic-Type Reactions and Part 3 Contaminants of Health or Environmental Concern.

NOI2005-01, List of Pest Control Product Formulants and Contaminants of Health or Environmental Concern under the New Pest Control Products Act.

The nature of the residues in plants and animals is adequately understood. The residue definition for enforcement is fluopyram in plant products and fluopyram including the metabolite fluopyram-benzamide in animal matrices. The approved uses of fluopyram on watermelon, wine grapes, strawberries, dry beans, dry chickpeas, dry lentils, peanuts, apples, potatoes, cherries and almonds does not constitute an unacceptable acute or chronic dietary risk (food and drinking water) to any segment of the population, including infants, children, adults and seniors. Sufficient crop residue data have been reviewed to recommend MRLs (see Table 3.5.1).

Mixers, loaders and applicators handling products containing fluopyram and workers re-entering treated areas are not expected to be exposed to levels of fluopyram that will result in risks of concern when the products are used according to label directions. The personal protective equipment on the product label is adequate to protect workers.

Residential exposure to individuals contacting treated fruits or foliage is not expected to result in risks of concern when products containing fluopyram are used according to label directions.

7.2 Environmental Risk

Fluopyram is persistent in soils and has a potential for long-term accumulation and residue carryover to the following crop season. Fluopyram is stable to hydrolysis, photolysis, aerobic and anaerobic biotransformation in soils. It does not form major transformation products in soils under Canadian field use conditions. Fluopyram is moderately mobile in soils and has a potential to leach and contaminate the groundwater depending on the soil type and location. Fluopyram has a low potential for bioconcentration/bioaccumulation in organisms.

Fluopyram is persistent in aquatic systems under aerobic and anaerobic conditions. Photolysis is not an important route of transformation in the aquatic environment. It does not form major transformation products in the water or sediment phases. Several minor transformation products were detected in natural water of which one was identified as fluopyram-lactam.

Fluopyram has a low potential for volatilization and, therefore, not expected to result in long range transport in the atmosphere.

Fluopyram presents a negligible risk to soil organisms, bees, beneficial arthropods, freshwater and marine fish, invertebrates, algae and aquatic plants. Fluopyram, however, may pose a risk to non-target terrestrial plants from spray drift (Luna Privilege only), and to amphibians due to runoff and spray drift. In order to minimize the potential risk, no-spray buffer zones between the treated area and downwind sensitive terrestrial and aquatic habitats are required. A bird toxicity label statement is also required as a precaution.

7.3 Value

The information submitted to register Luna Privilege, Luna Tranquility Fungicide, and Propulse Fungicide adequately demonstrated the value of the products in the management of a broad spectrum of foliar diseases and other fungal pathogens on various vegetable and fruit crops.

8.0 Regulatory Decision

Health Canada's PMRA, under the authority of the *Pest Control Products Act* and Regulations, has granted a conditional registration for the sale and use of the technical active, Fluopyram Technical Fungicide and end-use products, Luna Privilege containing the technical grade active ingredient fluopyram, Luna Tranquility Fungicide containing the technical grade active ingredients fluopyram and pyrimethanil and Propulse Fungicide containing the technical grade active ingredients fluopyram and prothioconazole to control various fungal diseases on various horticultural and field crops.

An evaluation of available scientific information found that, under the approved conditions of use, the products have value and do not present an unacceptable risk to human health or the environment.

Although the risks and value have been found acceptable when all risk-reduction measures are followed, as a condition of these registrations, additional scientific information (listed below) is being requested from the applicant as a result of this evaluation. For more details, refer to the Section 12 Notice associated with these conditional registrations.

NOTE: The PMRA will publish a consultation document at the time when there is a proposed decision on applications to convert these conditional registrations to full registrations or on applications to renew the conditional registrations, whichever occurs first.

Human Health

- DACO 4.3.1 Short term mode of action studies addressing the observed tumours. The goal of these studies is to further inform the two proposed cancer modes of action.
- DACO 7.2.3 (Inter-Laboratory Analytical Methodology Validation) An independent laboratory validation of Method GM-001-P07-01 for the determination of fluopyram residues in plant matrices is required to fulfill the data requirement for an acceptable enforcement method in plant matrices.
- DACO 7.4.4 (Field Accumulation Studies) A full set of field rotational crop data are required for canola, soybean and cereals (wheat, barley and corn, both field and sweet).

Value

- One field trial to confirm the efficacy of Luna Privilege against powdery mildew on standard sized cherry trees.
- One field trial to confirm efficacy of Luna Privilege against late leaf spot on peanuts.

List of Abbreviations

μg	micrograms
AB	Alberta
AD	administered dose
ADD	absorbed daily dose
ADI	acceptable daily intake
A:G	albumin/globulin
a.i.	active ingredient
ALAT	alanine aminotransferase
ALK	alkaline phosphatase
AR	applied radioactivity
ARfD	acute reference dose
ASAT	aspartate amino-transferase
BAF	bioaccumulation factor
BC	British Columbia
BCF	bioconcentration factor
BROD	benzyloxyresorufin O-deethylation
BW/bw	body weight
bwg	body weight gain
CAF	composite assessment factor
CAS	Chemical Abstracts Service
cm	centimetres
cm ²	centimetres square
cm ³	cubic centimetres
CO ₂	carbon dioxide
d d	day(s)
DFOP	Double-First-Order in Parallel
DFR	dislodgeable foliar residue
DT ₅₀	dissipation time 50% (the time required to observe a 50% decline in
	concentration)
DT ₇₅	dissipation time 75% (the time required to observe a 75% decline in concentration)
DT ₉₀	dissipation time 90% (the time required to observe a 90% decline in concentration)
dw	dry weight
EC ₀₅	effective concentration on 5% of the population
EC ₂₅	effective concentration on 25% of the population
EC ₅₀	effective concentration on 50% of the population
EDE	estimated daily exposure
EEC	estimated environmental exposure concentration
ER50	effective rate on 50% of the population
EROD	7-ethoxyresorufin <i>O</i> -deethylation
\mathbf{F}_{1}	first generation
F_2	second generation
FC	food consumption
FIR	food ingestion rate

fw fresh weight

g gram

GGT gamma glutamyltransferase

GIT gastrointestinal tract

h hour(s) ha hectare(s)

HAFT highest average field trial HDPE high-density polyethylene

HPLC high performance liquid chromatography

HPLC-MS/MS high performance liquid chromatography with tandem mass spectrometry

IBC intermediate bulk container IPM integrated pest management

IUPAC International Union of Pure and Applied Chemistry

K_d soil-water partition coefficient

kg kilogram

K_{oc} organic-carbon partition coefficient K_{ow} octanol-water partition coefficient

L litre

LADD lifetime average daily dose LC₅₀ lethal concentration 50%

LD low dose

LD₅₀ lethal dose 50%

LOAEC lowest observed adverse effect concentration

LOAEL lowest observed adverse effect level

LOC level of concern
LOQ limit of quantitation
LPM litre per minute
LR₅₀ lethal rate 50%
m metre(s)

m metre(s)
m³ cubic metre

MAS maximum average score

MB Manitoba mg milligram

MIS maximum irritation score

mL millilitre

M/L/A mixer/loader/applicator

MOA mode of action MOE margin of exposure

mol mole

MRL maximum residue limit
MTD maximum tolerated dose

N/A not applicable N/R not required

NAFTA North American Free Trade Agreement

NC not classified nanometre(s)

NOAEC no observed adverse effect concentration

NOAEL no observed adverse effect level

NOEC no observed effect concentration

NR not reported NS Nova Scotia

NZW New Zealand white

ON Ontario Pa Pascal

PBI plantback interval

PCA Fluopyram-pyridyl-carboxylic acid

PEI Prince Edward Island

PHED Pesticide Handlers Exposure Database

PHI preharvest interval dissociation constant

PMRA Pest Management Regulatory Agency

ppm parts per million

PROD pentoxyresorufin O-deethylation

PYO pick-your-own
O1* cancer potency factor

QC Quebec

RA risk assessment
RBC red blood cell
ROLD repeat oral low dose
RQ risk quotient

SFO single-first-order
SK Saskatchewan
SOHD single oral high dose
SOLD single oral low dose

STMdR supervised trial median residue STMR supervised trial mean residue

T3 tri-iodothyronine T4 thyroxine

TRR total radioactive residue
TSH thyroid stimulating hormone

TSMP Toxic Substances Management Policy

TWA time weighted average UDP uridine diphosphate US United States

UV ultraviolet wk week(s)

Appendix I Tables and Figures

Table 1 Residue Analysis

Matrix	Method ID	Analyte	Method Type	Limit of Quantitation		PMRA#
Soil	01068	fluopyram	HPLC- MS/MS	l μg/kg in soil		1599625
Soil	00973 01023	fluopyram AE C656948-benzamide (AE148815) AE C656948-7-hydroxy (BCS-AA-10065) AE C656948-PCA	HPLC- MS/MS	1 μg/kg in soil		1599622
Soil/ Sediment	GM-002-S07-01 GM-002-S07-04	fluopyram AE C656948-benzamide (AE148815) AE C656948-7-hydroxy (BCS-AA-10065) AE C656948-PCA	HPLC- MS/MS	1 μg/kg in soil and sedin	nent	1599627
Water	01051	fluopyram	HPLC- MS/MS	0.05 μg/L in drinking an	d surface water	1599623
Plant	GM-001-P07-01 enforcement method	fluopyram	HPLC- MS/MS	0.01 ppm	grape, strawberry, tomato	1599619
I —	benzamide, AE C656948-pyridy carboxylic acid, AE C656948-pyridy acetic acid, AE C656948-7-hydroxy	AE C656948-pyridyl- carboxylic acid, AE C656948-pyridyl-	HPLC- MS/MS	0.01 ppm for each analyte (except AE C656948-pyridyl- carboxylic acid on rape seed and AE C656948- methyl-sulfoxide on rape seed, wheat grain and lettuce)	lettuce head, rape seed, wheat grain, and orange	1599621
		sulfoxide		0.05 ppm for each analyte (except AE C656948-methyl- sulfoxide)	wheat straw	
	modification M001 to 00984	fluopyram, AE C656948- benzamide, AE C656948-pyridyl- carboxylic acid, AE C656948-pyridyl-acetic acid		0.01 ppm	processed commodities of apple, tomato, cabbage, grape, rape seed and strawberry	1599793, 1599737
Animal	01079 enforcement method	fluopyram and AE C656948-benzamide	HPLC- MS/MS	0.01 ppm	eggs, milk, fat, liver, kidney, muscle	1599626, 1599769
	Method 01061	Fluopy ram, AE C656948-benzamide, AE C656948-olefine (E- and Z- isomers)		0.01 ppm for fluopyram and AE C656948- benzamide; 0.02 ppm for calculated total residue of AE C656948-olefine (E- and Z- isomers)	eggs, milk, cream, fat, liver, kidney, muscle	1599624

Table 2 Toxicity Profile of Luna Privilege

(Effects are known or assumed to occur in both sexes unless otherwise noted; in such cases, sex-specific effects are separated by semi-colons)

Study Type/Animal	Study Results	PMRA#
Acute oral toxicity Wistar rats	Female LD ₅₀ > 2000 mg/kg bw Low toxicity	1599335
Acute dermal toxicity Wistar rats	$LD_{50} > 2000 \text{ mg/kg bw}$ Low toxicity	1599336
Acute inhalation toxicity (nose-only) Wistar rats	LC ₅₀ > 2.09 mg/L Low toxicity	1599337
Dermal irritation NZW rabbits	MAS = 0, MIS = 0 Non-irritating	1599338
Eye irritation NZW rabbits	MAS = 0, MIS = 5.3 Minimally irritating	1520933
Dermal sensitization (LLNA) CBA/J mouse	Non-sensitizer	1599340

Table 3 Toxicity Profile of Luna Tranquility Fungicide

(Effects are known or assumed to occur in both sexes unless otherwise noted; in such cases, sex-specific effects are separated by semi-colons)

Study Type/Animal	Study Results	PMRA#
Acute oral toxicity	Female LD ₅₀ > 5000 mg/kg bw	1670082
Sprague Dawley rats	Low toxicity	
Acute dermal toxicity	$LD_{50} > 2000 \text{ mg/kg bw}$	1670083
Wistar rats	Low toxicity	
Acute inhalation toxicity (nose-only)	$LC_{50} \ge 2.0 \text{ mg/L}$	1670084
Wistar rats	Low toxicity	
Dermal irritation	MAS = 0, $MIS = 0$	1670085
NZW rabbits	Non-irritating	
Eye irritation	MAS = 0, $MIS = 2$	1670086
NZW rabbits	Non-irritating	
Dermal sensitization (LLNA)	Non-sensitizer	1670087
CBA/J mouse		

Table 4 Toxicity Profile of Propulse Fungicide

(Effects are known or assumed to occur in both sexes unless otherwise noted; in such cases, sex-specific effects are separated by semi-colons)

Study Type/Animal	Study Results	PMRA#
Acute oral toxicity Sprague Dawley rats	Female LD ₅₀ : >5000 mg/kg bw Low toxicity	1670748
Acute dermal toxicity Sprague Dawley rats	LD ₅₀ : >5050 mg/kg bw Low toxicity	1670747
Acute inhalation toxicity (nose-only) Wistar rats	LC ₅₀ : ≥2.2 mg/L Low toxicity	1670744

Study Type/Animal	Study Results	PMRA#
Dermal irritation NZW rabbits	MAS = 0, MIS = 0 Non-irritating	1670745
Eye irritation NZW rabbits	MAS = 0, MIS = 0 Non-irritating	1670746
Dermal sensitization (Buehler) Guinea pigs	Non-sensitizer	1670749

Table 5 Toxicity Profile of Technical Fluopyram Fungicide

(Effects are known or assumed to occur in both sexes unless otherwise noted; in such cases, sex-specific effects are separated by semi-colons. Organ weight effects reflect both absolute organ weights and relative organ to bodyweights unless otherwise noted)

Study Type/Animal	Study Results	PMRA#
Metabolism/	Rate and extent of absorption and excretion: Fluopyram was rapidly and	1599513,
oxicokinetics (single and	effectively absorbed (93-98% of total recovered radioactivity; 3), as	1599517,
repeat dose, oral, gavage)	determined in a low dose (LD) bile-cannulation study. The AUC indicated	1599524,
Wistar rat	slightly higher systemic exposure for females than males in the single oral low	1599526,
	dose (SOLD; 5 mg/kg bw) tests and proportionality according to the dose.	1599529
	These findings were confirmed by quantitative whole body autoradiography.	
	Toxicokinetic data indicated major differences based on the part of the	
	molecule that had been radiolabelled. Time to maximum plasma	
	concentrations with the phenyl label were reached at 15 h in males and 11 h in	
	females (t _{max}) in SOLD animals, while that for repeat oral low dose (ROLD; 5	
	mg/kg bw/d - unradiolabelled for 14 d with radiolabelled fluopyram on day	
	15) animals was faster (0.8 h). The study with the pyridyl label produced	
	much shorter t _{max} values of 0.7 h and 3.3 h for SOLD males and females,	
	respectively. The C _{max} of single oral high dose of the phenyl label (SOHD;	
	250 mg/kg bw; phenyl) animals was between 35-42 h, suggesting delayed	
	absorption with increasing dose. There was evidence of an initial elimination	
	phase of 10-11 h, followed by a slower terminal elimination phase of 56-73 h	
	with a SOLD of the pyridyl label.	
	The majority of faecal and urinary excretion occurred within the first 72-96	
	hours; however, there was evidence of continuing excretion beyond 168 h as	
	evidenced by radioactive residues remaining in the carcass at sacrifice and the	
	confirmatory autoradiography results. Routes of excretion varied depending	
	on the location of the radiolabel. In males, faecal excretion accounted for	
	approx. 53% administered dose (AD), while urinary excretion ranged from 38-	
	45% AD. In females treated with the phenyl label, there were virtually equal	
	proportions of fluopyram excreted via the faeces and urine; in contrast, 39%	
	AD was faecal and 60% AD was urinary with the pyridyl label. Bile-	
	cannulated males showed total excretion of 90-100% AD, primarily due to	
	biliary excretion within the first 24 h, suggesting extensive enterohepatic	
	circulation. There were no significant levels of radiolabeled fluopyram in	
	expired air.	
	Distribution / target organ(s): Fluopyram was rapidly and widely distributed	
	in the body. The highest radioactive residues were observed in the liver,	
	kidney and Harderian gland, and in some studies, in the carcass, RBC, ovaries,	
	thyroid and adrenal glands. Total radioactivity remaining in the carcass was 2-	
	6% AD for the phenyl label and 0.3-0.5% AD for the pyridyl label. There was	
	some evidence of retention of fluopyram at 168 hours, particularly via the	
	renal route. No subsequent time-points were examined and thus the possibility	
	of bioaccumulation could not be excluded.	
	Toxicologically significant compound(s): Fluopyram was extensively	

Study Type/Animal	Study Results	PMRA#
Study Type/Animal	metabolized, with the ethyl linking group of the parent as the preferred site for metabolism, resulting in 7-hydroxy and 8-hydroxy metabolites. Further oxidation resulted in -enol, which was further conjugated to glucuronic acid. Hydroxylation of the phenyl ring resulted in -phenol and 7-OH-phenol metabolites. Elimination of water from compounds hydroxylated in the ethylene bridge resulted in fluopyram-Z-olefine and E-olefine metabolites (E-and Z-olefine can isomerize into each other). As the double bond of olefine may be a target for epoxidation and a dihydroxy-metabolite (which could result from hydrolysis of an epoxid by epoxid hydrolase) was observed, the olefine was considered to be of potential toxicological significance. All of the hydroxylated metabolites were conjugated primarily to glucuronic acid and to a lesser extent with sulfate. The cleavage of the molecule yielded label-specific metabolites (-benzamide; -pyridyl-acetic acid, -ethyl-diol, -pyridyl carboxylic acid) that represented the most abundant metabolites. This molecule was further metabolized via oxidation, hydroxylation and conjugation. The phenyl ring moiety was also conjugated with glutathione followed by further degradation to 7-OH-methyl-sulfone, -BA-methyl-sulfoxide and -BA-methyl-sulfone (phenyl label only). There were apparent sex differences in the quantity of metabolites generated. Fluopyram-7-hydroxy wand 7-OH-phenol metabolites were higher in males than females. Females showed higher amounts of 8-hydroxy and -benzamide than males. Low dose females excreted more of phenyl specific -benzamide and -benzoic acid than males. Females treated with the pyridyl label excreted more -pyridyl-acetic acid than males, while males excreted more -ethyl-diol metabolites than females. Parent accounted for 0.4/1.9% AD 3/\(\beta\) for the SOLD group and 10.5/16.7% AD 3/\(\beta\) for the SOHD group. Biliary	PMRA #
	metabolites were likely formed after first pass, with subsequent conjugation in GIT and subsequent excretion in facces. There were no significant differences	
	in metabolism between the doses, or between single and repeat dosing.	
Acute oral toxicity Wistar rats	Female LD ₅₀ : >2000 mg/kg bw Low Toxicity	1599564
Acute dermal toxicity Wistar rats	LD _{so} : >2000 mg/kg bw Low Toxicity	1599563
Acute inhalation toxicity (nose-only)	LC ₅₀ ; >5.1 mg/L air Low Toxicity	1599559
Wistar rats Skin irritation	MAS = 0, MIS = 0	1599561
NZW rabbits Eye Irritation	Non-irritating MAS = 1.8, MIS = 8. 7	1599558
NZW rabbits	Minimally irritating	
Skin Sensitization (LLNA) CBA/J mice	Non-sensitizer	1599573
4-weck dermal toxicity Wistar rat	Systemic NOAEL = 300 mg/kg bw/day Dermal NOAEL = 1000 mg/kg bw/day 1000 mg/kg bw/day: ↑ prothrombin time, ↑ cholesterol, ↑ liver weights, ↑ minimal centrilobular and mid-zonal hepatocellular hypertrophy No treatment-related dermal effects.	1599533
28-day dietary C57BL/6J mouse	Range-finding 24.7/31.1 mg/kg bw/day 3/9: ↑ liver weights, ↑ centrilobular hepatocellular hypertrophy	1599579

Study Type/Animal	Study Results	PMRA#
	162/197 mg/kg bw/day ♂/♀:	
	↑ liver weights, ↑ enlarged & dark livers, ↑ centrilobular hypertrophy, ↑ focal	
	necrosis in liver, ↑ single cell hepatocellular necrosis ♂; ↑ hypertrophy of	
	zona fasciculata in adrenals ♀	
	747/954 mg/kg bw/day ♂/♀ (exceeded MTD):	
	1 mortality due to intrathoracic hemorrhage, sacrificed Days 17-27, preceded	
	by severe clinical signs (↓ motor activity, hunched, piloerection, wasted appearance, cold to touch, laboured respiration, distended abdomen), marked bw loss, ↓ fc, ↑ pale pancreas, rounded borders in liver, dark & enlarged livers, reduced thymic size, distended abdomen, adrenal hypertrophy, vacuolation, degeneration/necrosis of zona fasciculate, perivascular & intra-alveolar hemorrhage of lungs, degeneration of pulmonary veins, erythroid extramedullary hematopoiesis in spleen, ↓ cellularity & focal hemorrhage, thyroid, centrilobular hypertrophy of hepatocytes, hepatocyte eosinophilia, bile duct/oval cell hyperplasia, focal necrosis, single cell necrosis; red liquid	
	thoracic cavity, centrilobular degeneration/necrosis &	
	Surviving females:	
	Distended abdomen, ↑ total cholesterol, total protein, ↑ ALAT, ↑ enlarged &	
	dark livers, ↑ hypertrophy of zona fasciculata in adrenals, ↑ centrilobular	
	hypertrophy in liver, ↑ single cell hepatocellular necrosis, ↑ focal necrosis in	
	liver, ↑ hepatocellular eosinophilia, ↑ bile duct/oval cell hyperplasia	
28-day dietary	Range-finding	1599574
Wistar rat		
	≥31.0/36.1 mg/kg bw/day ♂/♀:	
	↑ liver weights, ↑ enlarged, dark livers with prominent lobulation, ↑	
	centrilobular hepatocellular hypertrophy, ↑ pale kidneys, ↑ basophilic tubules,	
	hyaline droplets in proximal tubule, granular casts in medulla, ↑ P450, BROD,	
	PROD; ↑ thyroid weights, ↑ kidney weights ♂	
	254/263 mg/kg bw/day ♂/♀:	
	↓ bwg, ↑ total cholesterol, ↑ TG, ↑ follicular hypertrophy in thyroid, slight ↑	
	spleen weights; † colloidal depletion in thyroid, † platelets, † prothrombin	
	time, † size/cellularity follicles in spleen, † diffuse hypertrophy pituitary	
	basophils ∂, ↓ glucose; ↓ FC ♀	
28-day gavage	Supplemental	1599578
Beagle dog	750) mg/kg hw/dow	
	750 mg/kg bw/day: ↑ Soft/liquid/no feces, ↑ ALK, ↓ albumin and albumin globulin ratio, ↑ GGT,	
	↑ TG, ↑ enlarged livers ↑ liver weights, ↑ centrilobular-panlobular	
	hepatocellular hypertrophy, ↑ eosinophilic inclusions bodies; ↓ RBC, ↓	
	haemoglobin, ↓ hematocrit ♂	1599556
90-day dietary C57BL/6J mouse	NOAEL = $26.6/32.0$ mg/kg bw/day $3/9$	1399330
	188/216 mg/kg bw/day ♂/♀:	
	↑ ALAT, ↑ ALK, ↑ ASAT, ↓ albumin, ↑ adrenal weight, ↑ dark livers, ↑	
	focal necrosis in liver, ↑ cortical vacuolation in adrenals, ↓ cortical ceroid	
	pigment in adrenals	
90-day dietary	NOAEL = $12.5/14.6$ mg/kg bw/day $3/9$	1599557
Wistar rat		
	≥60.5/70.1 mg/kg bw/day ♂/♀:	
	bilirubin, ↑ TSH, ↑ T3, ↑ T4, pale kidneys, dark livers, ↑ prominent	
	lobulation in liver, positive cysts cortico-medullary junction, ↑ positive	

Study Type/Animal	Study Results	PMRA#
	cells/debris medulla of kidney, ↑ follicular cell hypertrophy in thyroid &; ↓	
	FC,↑ diffuse centrilobular hepatocellular hypertrophy, ↑ cholesterol ♀	
90-day dietary Dog	NOAEL = 28.5/32.9 mg/kg bw/day 3/9	1599555
	≥171/184 mg/kg bw/day ♂/♀:	
	bw, ↓ bwg; ↑ ALK, ↓ total bilirubin, ↓ albumin, ↓ A:G, ↓ total protein, ↑	
	liver weights, ↑ enlarged livers, ↑ hepatocellular hypertrophy &	
	intracytoplasmic eosinophilic droplets; ↓ FC, ↑ hepatocellular single cell	
	necrosis, ↑ incomplete maturation of prostate, zona glomerulosa vacuolation	
	in adrenals 3	
12-month dietary Beagle dog	NOAEL = 13.2/14.4 mg/kg bw/day ♂/♀	1599548
	67.6/66.1 mg/kg bw/day 3/2:	
	bwg wk 1, ↓ FC, ↑ ALK, ↑ GGT; ↑ diffuse hypertrophy of follicular	
	epithelium of thyroid, ↑ diffuse centrilobular hepatocellular hypertrophy &; ↑ thyroid weights ♀	
2-year dietary, chronic	Chronic toxicity:	1599635
toxicity / oncogenicity (combined)	NOAEL = $1.2/1.7$ mg/kg bw/day $3/9$	
Wistar rat	Liver carcinoma and adenoma ♀	
	52-week sacrifice:	
	6.0/8.6 mg/kg bw/day ♂/♀:	
	↑ liver centrilobular to panlobular hypertrophy, ↑ kidney weight, ↑ kidney,	
	histopathology (focal/multifocal chronic progressive nephropathy, hyaline	
	droplets in proximal tubules), ↑ cellular casts in urine, ↑ diffuse thyroid follicular cell hypertrophy ♂	
	Main group:	
	6.0/8.6 mg/kg bw/day ♂/♀:	
	↑ liver weight, ↑ liver histopathology (centrilobular to panlobular	
	hypertrophy, altered hepatocyte foci), Tenlarged kidney, Tkidney histopathology (chronic progressive nephropathy, focal/multifocal tubular	
	hyperplasia, focal/multifocal tubular dilatation, focal/multifocal tubular	
	hypertrophy), † thyroid follicular cell hypertrophy, † corneal opacity, corneal	
	oedema, nuclear opacity of lens, small retinal vessels 3 ; colloid alteration 9	
18 month dietary, chronic		1599632
toxicity / oncogenicity (combined)	NOAEL = $4.2/5.3$ mg/kg bw/day $3/2$	
C57BL/6J mouse	Thyroid follicular cell adenoma 3	
	52-week sacrifice:	
	20.9/26.8 mg/kg bw/day (♂/♀):	
	↑ liver weight; ↑ enlarged liver, ↑ focal/multifocal thyroid follicular cell	
	hyperplasia 3	
	Main group:	
	20.9/26.8 mg/kg bw/day (3/2):	
	↑ liver weight, ↑ diffuse centrilobular to panlobular hypertrophy, ↑	
	focal/multifocal thyroid follicular cell hyperplasia; ↑ focal/multifocal	
	hepatocellular single cell necrosis, ↑ platelets ♂; ↑ enlarged liver ♀	

Study Type/Animal	Study Results	PMRA#
One-generation dietary	Supplemental	1599823
range-finding)		
Wistar rat Supplemental	Parental effects	
	≥49.6/57.7 mg/kg bw/day ♂/♀:	
	↑ liver weights; ↑ kidney weight ♂	
	102.1/118.2 mg/kg bw/day ♂/♀:	
	thymus weight; ↓ premating bwg ♀	
Multi-generation dietary	Parental	1599824
Wistar rat	NOAEL = $13.9/16.8 \text{ mg/kg bw/day } 3/2$	
	Reproductive	
	NOAEL = $82.4/95.6$ mg/kg bw/day $3/9$	
	Offspring	
	NOAEL = $13.9/16.8 \text{ mg/kg bw/day} $	
	Parental effects	
	82.4/95.6 mg/kg bw/day 3/9:	
	↑ centrilobular hepatocellular hypertrophy; ↑ protein droplet nephropathy and	
	lymphocytic infiltration, \(\frac{1}{2}\) cytoplasmic vacuolization in adrenals \(F_1, \frac{1}{2}\) kidney	
	weights $F_0 \& F_1 \& f_2 \Leftrightarrow f_3 \Leftrightarrow f_4 \& f_5 \Leftrightarrow f_6 \& f_6 \Leftrightarrow f_6 \& f_6 \Leftrightarrow f_6 \& f_6 \Leftrightarrow f_6 \& f_6 \Leftrightarrow f_6 \Leftrightarrow f_6 \& f_6 \Leftrightarrow f_$	
	\uparrow cholesterol F_1 , \uparrow WBC F_1 , \uparrow monocyte absolute cell counts F_1 , \uparrow liver	
	weights $F_0 \& F_1$, \downarrow splcen weights $F_0 \& F_1$, \uparrow alveolar macrophages in lungs	
	$F_1 \subsetneq$	
	Offspring effects	
	82.4/95.6 mg/kg bw/day ♂/♀:	
	\downarrow bw F_1 & F_2 , \downarrow bwg F_1 & F_2 , \downarrow spleen and thymus weights F_2	1599610
Developmental toxicity Sprague Dawley rat	Maternal NOAEL = 30 mg/kg bw/day	1399010
Sprague Dawiey rat	TOTAL SO MIGHE OWNERS	
	Developmental	
	NOAEL = 150 mg/kg bw/day	
	Maternal effects:	
	≥150 mg/kg bw/day:	
	↓ bwg, ↓ corrected bwg to gravid uterine weight, ↓ FC, ↑ abs. liver weight, ↑	
	centrilobular hepatocellular hypertrophy	
	Developmental effects:	
	450 mg/kg bw/day:	
	↓ fetal weights, ↑ thymic remnant present, ↑ skeletal variations	
	No evidence of teratogenicity	
Developmental toxicity	Maternal	1599571
NZW rabbit	NOAEL = 25 mg/kg bw/day	
	D 1	
	Developmental effects	
	NOAEL = 25 mg/kg bw/day	
	Maternal effects:	
	75 mg/kg bw/day:	
	bw, ↓ bwg, ↓ corrected bwg to gravid uterine weight, ↓ FC	
	Developmental effects:	
	75 mg/kg bw/day:	
	↓ fetal weights, ↑ runts (bw < 28.0 g)	

Study Type/Animal	Study Results	PMRA#
Acute Neurotoxicity Wistar rat	Main study: NOAEL = 125 mg/kg bw/not established ♂/♀ Supplemental study (female only): Female NOAEL = 50 mg/kg bw Main study: ≥125 mg/kg bw: ↓ session motor activity, ↓ session locomotor activity ♀ ≥ 500 mg/kg bw: ↓ session motor activity, ↓ session locomotor activity ♂; ↓ body temperature, ↓ vocalization during removal ♀ Supplemental: 100 mg/kg bw:	1599618
Subchronic Neurotoxicity Wistar rat	\$\delta\$ session motor activity, \$\delta\$ session locomotor activity \$\times\$ Systemic toxicity: NOAEL = 33.2/41.2 mg/kg bw/day \$\delta / \times\$ 164.2/197.1 mg/kg bw/day \$\delta / \times\$: \$\delta\$ bw, \$\delta\$ bwg, \$\delta\$ cholesterol, \$\delta\$ bilateral retinal degeneration; \$\delta\$ FC \$\delta\$; \$\delta\$ TG, \$\delta\$ thyroid weights \$\times\$	1599534
Gene mutations in bacteria in vitro	Negative	1599580
Gene mutations in bacteria in vitro	Negative	1599553
In vitro mammalian clastogenicity Chromosome aberrations	Negative	1599552
In vitro mammalian cell assay V79/HPRT forward mutation	Negative	1229493
In vivo cytogenetics Micronucleus assay	Negative	1229494
3-day toxicity study in male C57BL/6J mouse – pharmacokinetic investigations of the clearance of intravenous (iv)-administered ¹²⁵ I-thyroxine	Non-guideline Whole blood thyroxine levels were lower in fluopyram-treated males at all time-points compared to controls. Similar effects were observed in PB-treated males, although the decreases from controls were marginally less and there was some evidence of recovery at 24 h.	1654272
3-day toxicity study in male C57BL/6J mice - QPCR investigations of gene transcripts in the liver	Non-guideline Fluopyram (300 mg/kg bw/day): ↑ liver weight, ↑ expression of the following genes: Cyp1a, Cyp2b, Cyp 3a, Sult1a1, Sult 2a2, Suln, Ugt1a1, Ugt2b1, Ugt2b5 PB (80 mg/kg bw): Reduced motor activity ↑ liver weight, ↑ expression of the following genes: Cyp2b, Cyp 3a, Sult1a1, Sult 2a2, Suln, Ugt1a1, Ugt2b1, Ugt2b5	1654273

Study Type/Animal	Study Results	PMRA#
Study Type/Animal 3-day, 14-day toxicity in male C57BL/6J mice (dietary) – hepatotoxicity and thyroid hormones fluopyram vs. phenobarbital	Non-guideline 3 day exposure 308 mg/kg bw/day: ↓ FC, ↓ T4, ↑ TSH, ↑ liver weight, ↑ enlarged livers, ↑ centrilobular to panlobular hepatocellular hypertrophy (diffuse; graded minimal-slight), ↑ number of mitoses present, ↑ hepatocellular single cell necrosis, ↑ total P450, EROD, PROD, BROD Phenobarbital at 80 mg/kg bw/day: bw loss, ↓ FC, ↓ T4, ↓ T3, ↑ rel. liver weight, ↑ enlarged and dark livers, ↑ centrilobular to panlobular hepatocellular hypertrophy (diffuse; graded minimal-slight), ↑ number of mitoses present, ↑ total P450, EROD, PROD, BROD 14 day exposure 314 mg/kg bw/day: ↓ FC, ↓ T4, ↑ TSH, ↑ liver weight, ↑ enlarged and dark livers, ↑ centrilobular to panlobular hepatocellular hypertrophy (diffuse; graded slight-moderate), ↑ hepatocellular single cell necrosis, ↑ total P450, EROD, PROD, BROD	PMRA # 1599576, 1599803
	Phenobarbital at 80 mg/kg bw/day: bw loss, ↓ FC, ↓ T4, ↑ TSH, ↑ liver weight, ↑ enlarged and dark livers, ↑ centrilobular to panlobular hepatocellular hypertrophy (diffuse; graded slightmoderate), ↑ hepatocellular single cell necrosis, ↑ total P450, EROD, PROD, BROD	1500741
7-day toxicity in female Wistar rats (dictary) fluopyram vs. phenobarbital	Non-guideline 193 mg/kg bw/day: ↓ FC, ↑ liver weight, ↑ enlarged and dark livers, ↑ centrilobular to panlobular hepatocellular hypertrophy (diffuse; graded minimal-slight), ↓ diffuse mainly periportal hepatocellular vacuolation, ↑ BrdU labelling index in centrilobular and periportal zones of liver, ↑ total P450, EROD, PROD, BROD, UDPGT Phenobarbital at 80 mg/kg bw/day: Reduced motor activity, ↓ bw, ↓ bwg, ↑ liver weight, ↑ enlarged and dark livers, ↑ centrilobular to panlobular hepatocellular hypertrophy (diffuse; graded minimal-slight), ↑ hepatocellular necrotic focus, ↑ BrdU labelling index in centrilobular and periportal zones of liver, ↑ total P450, EROD, PROD, BROD, UDPGT	1599741, 1599802
In vitro studies with hog thyroid microsomes on the potential interactions with thyroid peroxidase- catalyzed reactions	Non-guideline Fluopyram does not affect thyroid hormone synthesis at the level of TPO under the study conditions tested. The effects of fluopyram metabolites were not studied.	1599551
	656948-pyridyl-carboxylic acid (AE 657188)	
Acute oral toxicity	Female LD ₅₀ :>2000 mg/kg bw	1599809
Wistar rats	Low Toxicity	1500712
28-day dietary Sprague Dawley rat	NOAEL = 1574/162 mg/kg bw/day ♂/♀ 1581 mg/kg bw/day ♀: ↓ bwg, FC	1599612

Study Type/Animal	Study Results	PMRA#
	Negative	1599630
In vitro mammalian cell	Precipitation observed at ≥4000 μg/mL in the absence of S9 mix and at 5000 μg/mL in the presence of S9 mix (data were not interpretable). Negative	1599613
Chromosome aberrations	Negative	1599611

Table 6 Toxicology Endpoints for Use in Health Risk Assessment for Fluopyram

Exposure Scenario	Study	Point of Departure and Endpoint	CAF ¹ or Target MOE
Acute dietary general population	Rat acute neurotoxicity study	NOAEL = 50 mg/kg bw Reduced motor and locomotor activity	100
	Acute reference dose = 0.5 mg/	kg bw	
Repeated dietary	Rat chronic toxicity/carcinogenicity study	NOAEL = 1.2 mg/kg bw/day Numerous effects, primarily in liver, kidney, thyroid and eye	100
	Acceptable daily intake = 0.012	mg/kg bw/day	
Short- and intermediate term dermal ²	Rat 28 day dermal toxicity study	NOAEL = 300 mg/kg bw/day Clinical chemistry and liver effects	100
Long-term dermal ²	Rat chronic toxicity/carcinogenicity study	NOAEL = 1.2 mg/kg bw/day Numerous effects, primarily in liver, kidney, thyroid and eye	100
Short- and intermediate- term inhalation ³	Rat 90 day oral toxicity study	NOAEL = 12.5 mg/kg bw/day Numerous effects	100
Long-term inhalation ³	Rat chronic toxicity/carcinogenicity study	NOAEL = 1.2 mg/kg bw/day Numerous effects, primarily in liver, kidney, thyroid and eye	100
Pick-your-own and residential ornamental oral	Rat acute neurotoxicity study	NOAEL = 50 mg/kg bw Reduced motor and locomotor activity	100
Pick-your-own and residential dermal	Rat 28 day dermal toxicity study	NOAEL = 300 mg/kg bw/day Clinical chemistry and liver effects	100
Cancer	O_1 * set at 1.72 × 10 ⁻² (mg/kg bw	/day) ⁻¹	

CAF (composite assessment factor) refers to a total of uncertainty and *Pest Control Products Act* factors for dietary assessments; MOE refers to a target MOE for occupational and residential assessments

Since an oral NOAEL was selected, a dermal absorption factor was used in a route-to-route extrapolation
 Since an oral NOAEL was selected, an inhalation absorption factor was used in route-to-route extrapolation.

Table 7 Exposure Estimates for Mixers/Loaders/Applicators

Scenario	Area Treated	,	Unit Exposu ag/kg a.i. han				
	Non-Cancer	Cancer	Dermal	Inhalation	Combined		
Groundboom- Fruits and Vegetables Farmer/Custom	26	12	84.12	2.56	86.68		
Groundboom Farmer – large field crops	107	60	84.12	2.56	86.68		
Groundboom Custom – large field crops	360	240	84.12	2.56	86.68		
Airblast	20	7	879.38	7.4	886.78		
Drip Application Mix/Load	26	12	51.14	1.6	52.74		
Aerial Mix/Load	400	318	51.14	1.6	52.74		
Aerial Applicator	400	318	9.66	0.07	9.73		

Table 8 Non-Cancer Exposure and Risk Estimates for Mixer/Loader/Applicators Handling Fluopyram

Crop	Application Equipment	Maximum Rate kg a.i./ha	Exposure	Inhalation Exposure mg/kg bw/day	Dermal MOE ^a	Inhalation MOE ^a
Watermelon	groundboom	0.25	0.00781	0.000238	38407	52584
Wine Grapes	airblast	0.25	0.0628	0.000529	4776	23649
Dry beans	groundboom farmer	0.15	0.0193	0.000587	15554	21296
	groundboom custom	0.15	0.0649	0.00198	4623	6330
Peanuts	groundboom	0.25	0.0321	0.000978	9332	12777
Apples	airblast	0.15	0.0377	0.000317	7960	39414
Potatoes	groundboom farmer	0.15	0.0193	0.000587	15554	21296
	groundboom custom	0.15	0.0649	0.00198	4623	6330
	aerial M/L	0.15	0.0438	0.00137	6844	9115
	aerial applicator	0.15	0.00828	0,00006	36232	208333
Strawberries	drip irrigation	0.25	0.004749	0.000149	63175	84135
Cherries	airblast	0.125	0.0314	0.000264	9552	47297
Almonds (tree nuts)	airblast	0.25	0.0628	0.000529	4776	23649

a Target MOE = 100

Table 9 Cancer Exposure and Risk Estimates for Mixer/Loader/Applicators Handling Fluopyram

Crop	Application Equipment	Maximum Appln Rate kg a.i./ha	Maximum Number of Apps/season	Dermal Exposure mg/kg bw/day	Inhalation Exposure mg/kg bw/day	LADD mg/kg bw/day	Cancer Risk
Watermelon	groundboom	0.25	2	0.000252	0.00011	1.06E-06	1.82E-08
Wine Grapes	airblast	0.25	2	0.001539	0.000185	5.04E-06	8.67E-08
Dry beans	groundboom farmer	0.15	2	0.000757	0.000329	3.17E-06	5.46E-08
	groundboom custom	0.15	60	0.003028	0.001317	3.81E-04	6.55E-06
Peanuts	groundboom	0.25	2	0.001262	0.000549	5.29E-06	9.10E-08
Apples	airblast	0.15	3	0.000923	0.000111	4.53E-06	7.80E-08
Potatoes	groundboom farmer	0.15	2	0.000757	0.000329	3.17E-06	5.46E-08
	groundboom custom	0.15	60	0.003028	0.001317	3.81E-04	6.55E-06
	aerial M/L	0.15	2	0.002439	0.00109	3.09E-04	5.32E-06
	aerial applicator	0.15	2	0.000461	4.77E-05	4.46E-05	7.67E-07
Strawberries	drip irrigation	0.25	2	0.000153	6.86E-05	6.49E-07	1.12E-08
Cherries	airblast	0.125	3	0.000769	9.25E-05	3.78E-06	6.50E-08
Almonds (tree nuts)		0.25	2	0.001539	0.000185	5.04E-06	8.67E-08

Table 10 Non-Cancer Postapplication Exposure and Risk Estimates for Fluopyram

Crop	Reentry Activity	Maximum Appln Rate kg a.i./ha	Max Number of Apps/season	Transfer Coefficient cm²/h	DFR Value µg/cm²	Dermal Exposure mg/kg bw/day	Dermal MOE ^a
Watermelon	hand harvesting, leaf pulling, hand pruning, thinning, turning	0.25	2	2500	0.7391	0.211	1421
Wine Grapes	hand harvesting, training, thinning, hand pruning, tying, leaf pulling	0.25	2	8500	0.7391	0.718	418
	scouting, irrigation	0.15	2	1500	0.4435	0.076	3946
Dry beans	hand harvesting (green peas)	0.15	2	2500	0.4435	0.127	2368
Peanuts	scouting, irrigation	0.25	2	1500	0.6144	0.105	2848
Apples	thinning	0.15	3	3000	0.5121	0.176	1709
Potatoes	scouting, irrigation	0.15	2	1500	0.4435	0.076	3946
	hand harvest (sweet potatoes)	0.15	2	2500	0.4435	0.127	2368
Strawberries	hand harvesting, thinning, hand pruning, tying, training	0.25	2	1500	0.7952	0.136	2201
Cherries	thinning	0.125	3	3000	0.4268	0.146	2050
Almonds (tree nuts)	harvesting	0.25	2	200	0.6144	0.014	21362

a Target MOE = 100

Table 11 Cancer Postapplication Exposure and Risk Estimates for Fluopyram

Crop	Reentry Activity	Maximum Appln Rate kg a.i./ha	Exposure Frequency days/year	30 day TWA DFR Value µg/cm ²	ADD mg/kg bw/day	LADD	Cancer Risk
Watermelon	hand harvesting, leaf pulling, hand pruning, thinning, turning	0.25	30	0.317	0.00634	0.000278	4.8E-06
Wine Grapes	hand harvesting, training, thinning, hand pruning, tying, leaf pulling	0.25	30	0.317	0.02156	0.000945	1.6E-05
Dry beans	scouting, irrigation	0.15	30	0.190	0.00228	0.000100	1.7E-06
	hand harvesting (green peas)	0.15	30	0.190	0.00380	0.000167	2.9E-06
Peanuts	scouting, irrigation	0.25	30	0.295	0.00354	0.000155	2.7E-06
Apples	thinning	0.15	30	0.273	0.00655	0.000287	4.9E-06
Potatoes	scouting, irrigation	0.15	30	0.190	0.00228	0.000100	1.7E-06
	hand harvest (sweet potatoes)	0.15	30	0.190	0.00380	0.000167	2.9E-06
Strawberries	hand harvesting, thinning, hand pruning, tying, training	0.25	30	0.322	0.00386	0.000169	2.9E-06
Cherries	thinning	0.125	30	0.227	0.00546	0.000239	4.1E-06
Almonds (tree nuts)	harvesting	0.25	30	0.295	0.00047	0.000021	3.6E-07

Major Groundwater and Surface Water Model Inputs for Level 1, Level 2 Table 12 and Level 2 Restricted Application Assessments

Type of Input	Parameter	Value
Application information:	Crop(s) to be treated	grapes, apples, water melon, wine grapes, dry beans, peanut, potato, cherry, and tree nuts
	Maximum allowable application rate per year (g a.i./ha)	500
	Maximum rate for each application (g a.i./ha)	250
	Maximum number of applications per year	2
	Minimum interval between applications (days)	7
	Method of application	Airblast
Application information:	Crops to be treated	grapes potato (drinking water only)
Level 2 (Dugout	Maximum allowable application rate per year (g a.i/ha)	500
only)	Maximum rate each application (g a.i./ha)	250
	Maximum number of applications per year	2
	Minimum interval between applications (days)	1) 7-days 2) 14-days (drinking water)
	Method of application	Ground
Environmental fate	Hydrolysis half-life at pH 7 (days)	Stable
characteristics	Photolysis half-life in water (days)	Stable
	Adsorption K_{∞} (mL/g)	284 (20 th percentile of five K_{∞} values for fluopyram)
	Aerobic soil biotransformation half-life (days)	654 for Level 1 and Level 2 (80 th percentile of half-life values; values for the two labels were averaged) 533 at Level 2 restricted application (80 th percentile of fitted lognormal distribution; values for the two labels were averaged)
	Aerobic aquatic biotransformation half-life (days)	1330 (longest of two half-lives; values for the two labels were averaged)
	Anaerobic aquatic biotransformation half-life (days)	1495 (single half-life; values for the two labels were averaged)

Level 1 and Level 2 Estimated Environmental Concentrations of Fluopyram Table 13 in Potential Drinking Water Sources

	Groundw	vater EEC	Su	rface Water	Vater EEC (µg a.i./L)			
Compound	(µg а	a.i./L)	Rese	ervoir	Du	ugout		
	Daily ¹	Yearly ²	Daily ³	Yearly ⁴	Daily ³	Yearly4		
Fluopyram, Level 1	106	104	26	7.8	236	231		
Fluopyram, Level 2	N/A	N/A	N/A	N/A	185	181		

Notes:

- 90th percentile of daily average concentrations
- 90th percentile of yearly average concentrations 90th percentile of yearly peak concentrations 90th percentile of yearly average concentrations 90th percentile of yearly average concentrations

N/A = not applicable

Table 14 Level 2 Additional Modelling - Restricted Application Estimated
Environmental Concentrations of Fluopyram in Potential Drinking Water
Sources

Use pattern	Groundw	ater EEC	Sur	EEC (µg a	C (µg a.i./L)	
	(µg а	.i./L)	Rese	ervoir	Du	gout
	Daily ³	Yearly4	Daily1	Yearly ²	Daily1	Yearly ²
Apply one year only	15	15	17	NR	13	12
Apply two years only	28	28	26	NR	22	21
Apply three years only	40	40	26	NR	26	25

Notes: 1 90th percentile of yearly peak concentrations,

2 90th percentile of yearly average concentrations

3 90th percentile of maximum daily concentration for each of twelve starting years

4 90th percentile of maximum yearly concentration for each of twelve starting years

N/A For level 2, the reservoir and groundwater were not modelled.

N/R The yearly values were not reported as EECs would decline rapidly after the number of years of application, whether 1, 2 or 3 years. In other words, there is only one peak for each year of application and concentration declines to almost zero in subsequent years. Therefore, the 90th percentile of yearly averages calculated for these restricted years

Table 15 Groundwater EECs (µg/L) Averaged over Five Time Periods*

N years	AB_ North	AB_S_ irr	BC_F irr	BC_O_ irr	SK_ Rgina	MB_ Wnpeg	ON_ Essex	ON_ Niaga	QC_ Yamsk	PEI_ Char	NS_ Fundy
Highest	daily EF	EC									
1	7	14.7	8	8.6	0.5	2.5	6.5	4.4	0.8	6.2	10.6
2	13.9	28.3	14.5	16.4	0.8	4.7	13.5	8.9	1.6	11.9	20.1
3	20.7	40.4	19.1	23.3	1.1	6.8	20.1	13.2	2.4	17.5	28.3
Highest	one year	average	value								
1	6.9	14.7	7.8	8.5	0.5	2.5	6.4	4.4	0.8	6.1	10.5
2	13.6	28.2	14.1	16.4	0.7	4.7	13.2	8.8	1.6	11.8	19.8
3	20.4	40.2	18.6	23.2	1	6.8	20	13.1	2.4	17.2	27.7
Five yea	r averag	e EEC		-							
1	6.7	12.1	4.5	7.9	0.4	2.4	5.8	3.9	0.8	5.4	8.6
2	13.4	23.9	8.8	15.2	0.7	4.6	11.8	7.8	1.5	10.5	17
3	20	34.8	12.7	21.5	0.9	6.6	17.4	11.9	2.4	15.3	24.2
Ten year	averag	e EEC									
1	6.6	8.1	2.3	6.2	0.3	2.3	4.4	3.2	0.7	3.9	6
2	13	16.3	4.6	11.9	0.6	4.3	9.1	6.4	1.4	7.7	11.7
3	19.4	24.4	6.9	17.4	0.8	6.3	13.6	9.7	2.2	11.2	17
Twenty	year ave	rage EE	C								
1	5.6	4.4	1.2	3.9	0.3	1.9	2.6	2.1	0.6	2.2	3.2
2	11.2	8.9	2.4	7.4	0.4	3.5	5.4	4.3	1.2	4.4	6.3
3	16.6	13.4	3.6	10.7	0.5	5	8.1	6.4	1.8	6.5	9.3
Seventy	year ave	erage EE	C								
1	1.95	1.19	0.33	1.07	0.08	0.65	0.72	0.62	0.19	0.61	0.88
2	3.86	2.44	0.66	2.05	0.14	1.22	1.48	1.24	0.37	1.21	1.74
3	5.77	3.68	0.99	2.96	0.18	1.76	2.23	1.87	0.57	1.78	2.55

*daily, one year, five years, ten years and twenty years and seventy years - 500 g a.i./ha (two applications at 250 g a.i./ha per year)

Table 16 Number of Days When EECs Exceed 2 μg/L for All 11 Groundwater Scenarios, Assuming Applications over One, Two or Three Years

N years	AB_ North	AB_S_ irr	BC_F_i	BC_O_ irr	SK_ Rgina	MB_ Wnpeg	ON_ Essex	ON_ Niaga	QC_ Yamsk	PEI_ Char	NS_ Fundy
1	9297	6671	1435	4379	0	3384	3630	3332	0	3155	3190
2	11739	8063	1879	5465	0	6404	4768	5109	0	4083	3865
3	12997	8964	2263	6179	0	8114	5198	5877	2928	4696	4323

Table 17 Groundwater EECs (µg/L)* Averaged over Seventy Years

N years*	AB_ North		BC_F_ irr	BC_O_ irr				ON_ Niaga	QC_ Yamsk	PEI_ Char	NS_ Fundy
3	4.61	2.93	0.79	2.37	0.14	1.41	1.78	1.50	0.46	1.43	2.04
100	81	59	19	61	0.53	30	42	38	11	36	42

^{*}averaged over 70 years assuming three or 100 consecutive years of application of fluopyram at a reduced potato use rate of 400 g a.i./ha per year (two applications of 150 g a.i./ha plus one of 100 g a.i./ha at an interval of 7 days)

Table 18a Nature of the Residues in Plant Matrices

Nature of the Residue in	Grapes		PMRA# 1599785 and 1599786				
Radiolabeled Position		[phenyl-UL-14C] fluopyram and [pyridyl-2,6-14C] fluopyram					
Test Site	Plants were grown under natural sunlight and temperatures, except that a glass roof wa automatically closed at the beginning of rainfall.						
Treatment	Three foliar spray applications at 100, 200 and 200 g a.i./ha; intervals between applications were 42 and 49 days.						
Rate	504 g a.i./ha (phenyl) and	504 g a.i./ha (phenyl) and 498 g a.i./ha (pyridyl)					
End-use Product	Fluopyram 500 SC						
Preharvest interval	18 days						
Matrix	PHI (days)	[phenyl-14 TRRs (ppn		[pyridyl-14C] TRRs (ppm)			
Summer Cut	After second application	28.55		64.18			
Grapes	18	1.86		1.70			
Leaves	19	48.06		42.66			
Metabolites Identified	Major Metabolites (>10%	of the TRRs)	Minor Metabolites (<10% of the TRRs)				
[phenyl-14C]							
Summer Cut	Fluopyram		None				
Grapes	Fluopyram		AE C656948-benzamide AE C656948-7-hydroxy				
Leaves	Fluopyram		AE C656948-7-hydroxy AE C656948-8-hydroxy glucoside conjugate of AE C656948-7- hydroxy				
[pyridyl-14C]							
Summer Cut	Fluopyram		AE C656948-pyridyl-carboxylic acid AE C656948-7-hydroxy AE C656948-8-hydroxy glucoside conjugate of AE C656948-7- hydroxy				
Grapes	Fluopyram		AE C656948-pyridyl-carboxylic acid AE C656948-7-hydroxy				

Leaves	Fluopyram		AE C656948-pyridyl-carboxylic acid AE C656948-7-hydroxy AE C656948-8-hydroxy glucoside conjugate of AE C656948-7- hydroxy				
1.0% of the TRRs. The Hydroxylation of fluopy Conjugation of AE C65	main reactions involved ram leading to AE C656 6948-7-hydroxy	are: 6948-7-hydroxy and	of the metabolite AE C656948-8-				
		ng to AE C050948-t		AE C656948-carboxylic acid 9781 and 1599789			
Nature of the Residue Radiolabeled Position		luopyram and [pyrid					
Test Site	Plants were grown	under natural sunligl	nt and temperatu	ares, except that a glass roof w			
Treatment	Three foliar spray a	automatically closed at the beginning of rainfall. Three foliar spray applications at approximately 167 g a.i./ha; intervals between applications were 16 and 11 days.					
Data		nyl) and 505.7 g a.i./	(ha (nyridyl)				
Rate End-use Product	Fluopyram 500 SC	ny 1) and 202.1 g a.l.	in (pyridyr)				
Preharvest interval	51 days						
Matrix	PHI (days)	[phenyl-140 TRRs (ppm		[pyridyl-14C] TRRs (ppm)			
Potato tuber	51	0.008	1)	0.012			
Potato leaves	51	47.64		21.67			
Metabolites Identified		(>10% of the TRRs)	Minor Metab	olites (<10% of the TRRs)			
	iviajoi ivictatorines	(~1076 01 the 11kKs)	Ivillioi ivictao	ones (1070 of the 11dts)			
[phenyl-14C] Potato tuber	Fluopyram		AE C656948	henzamide			
Potato tuber	riuopyram		AE C656948-7-hydroxy				
Potato leaves	Fluopyram		AE C656948-benzamide				
Potato leaves	riuopyram		AE C656948-7-hydroxy				
[pyridyl-14C]			1112 0000710	· My date of			
Potato tuber	Fluopyram AE C656948-pyrid (49.8% of the TRR		AE C656948-pyridyl-carboxylic acid AE C656948-7-hydroxy				
Potato leaves	Fluopyram		AE C656948-pyridyl-carboxylic acid AE C656948-7-hydroxy				
The metabolic pathway	of fluopyram in potatoes	s consisted of hydrox	xylation of fluor	yram leading to AE C656948			
hydroxy, which is cleav	ed to AE C656948-benz	amide and AE C656	948-PCA.				
Nature of the Residue	in Beans		PMRA # 159	9779 and 1599787			
Radiolabeled Position	[phenyl-UL-14C] f	luopyram and [pyrid	yl-2,6-14C] fluo	ppyram			
Test Site	automatically close	Plants were grown under natural sunlight and temperatures, except that a glass roof wa automatically closed at the beginning of rainfall.					
Treatment	applications was 28	Two foliar spray applications at approximately 250 g a.i./ha; the interval between applications was 28 days.					
Rate		528 g a.i./ha (phenyl) and 519 g a.i./ha (pyridyl)					
End-use Product	Fluopyram 500 SC						
Preharvest interval	4 days for immatur	4 days for immature crops and 29 days f					
Matrix	PHI			[pyridyl-14C]			
	(days)	TRRs (ppm)	TRRs (ppm)			
Green bean	4	1.40		3.88			
Foliage	4	36.66		38.53			
Succulent bean	29	0.07		0.17			
Dry beans	29 + drying for 11			0.31			
Straw	29	16.55		19.02			
Metabolites Identified		(>10% of the TRRs)		olites (<10% of the TRRs)			

Green bean	Fluopyram	None			
Foliage	Fluopyram	AE C656948-benzamide			
		AE C656948-7-hydroxy			
		AE C656948-8-hydroxy			
		glucoside conjugate of AE C656948-7-			
		hydroxy			
Succulent beans	Fluopyram	AE C656948-7-hydroxy			
succulent beans	AE C656948-benzamide	AE C656948-8-hydroxy			
	(51.9% of the TRRs; 0.036 ppm)	glucoside conjugate of AE C656948-7-			
	(31.9% of the TRRS, 0.930 ppin)	hydroxy			
2 1	E1				
Dry beans	Fluopyram	AE C656948-7-hydroxy			
	AE C656948-benzamide	AE C656948-8-hydroxy			
	(64.0% of the TRRs; 0.077 ppm)	glucoside conjugate of AE C656948-7-			
		hydroxy			
Straw	Fluopyram	AE C656948-benzamide			
		AE C656948-7-hydroxy			
		AE C656948-8-hydroxy			
		glucoside conjugate of AE C656948-7-			
		hydroxy			
pyridyl-14C]					
Green bean	Fluopyram	None			
Foliage	Fluopyram	AE C656948-PCA			
· Ond Bo	1.00	AE C656948-7-hydroxy			
		AE C656948-8-hydroxy			
		glucoside conjugate of AE C656948-7-			
		hydroxy			
7 1 11	AE C656948-PAA				
Succulent beans		Fluopyram			
	(29.5% of the TRRs; 0.051 ppm)	AE C656948-7-hydroxy			
	AE C656948-PCA	AE C656948-8-hydroxy			
	(31.0% of the TRRs; 0.054 ppm)	glucoside conjugate of AE C656948-7-			
		hydroxy			
Dry beans	AE C656948-PAA	Fluopyram			
	(22.6% of the TRRs; 0.070 ppm)	AE C656948-7-hydroxy			
	AE C656948-PCA	AE C656948-8-hydroxy			
	(32.5% of the TRRs; 0.100 ppm)	glucoside conjugate of AE C656948-7-			
		hydroxy			
Straw	Fluopyram	AE C656948-PCA			
		AE C656948-PAA			
		AE C656948-7-hydroxy			
		AE C656948-8-hydroxy			
		glucoside conjugate of AE C656948-7-			
		hydroxy			
The main reactions of flu	opyram metabolism in the beans are:	inymony			
	ram leading to AE C656948-7-hydroxy at	nd AE C656048 & hydroxy			
		c acid, in one case conjugation of AE C656948-8-			
hydroxy with glycoside a		acid, in one case conjugation of AL C030748-8			
nydroxy with glycoside a	d active substance leading to AE C65604	8-benzamide and AE C656948-carboxylic acid			
Nature of the Residue in		PMRA# 1599782 and 1599790			
Radiolabeled Position	[phenyl-UL-14C] fluopyram and [py				
Test Site	Soil-less cultivation (stone wool subs	strate) in a greenhouse			
Treatment	Drip irrigation				
Rate		5 mg a.i./plant. Additionally, an experiment was			
conducted at an exaggerated rate (4x) of 20 mg a.i./plant.					
	conducted at an exaggerated rate (4x)	of 20 mg a.i./plant.			

Preharvest interval	Intermediate plant (4x experiment Mature peppers (1x experiment, 196 days) Mature peppers (4x experiment, 155 to 96 days) Rest of plant (1x experiment, 97 days)	both radio pyridyl rad	labels, three ti		
Matrix		enyl-14C		[pyridyl-14C]	
		Rs (ppm)		TRRs (ppm)	
Pepper Intermediate (4x)	33 6.2	37	,	18.24	
Mature peppers (1x)	55-96 0.0			0.060	
Mature peppers (4x)		t applicab	le	0.149	
Rest of plant (1x)	97 3.5		10	2.344	
Metabolites Identified	Major Metabolites (>10% of the		Minor Metabo	lites (<10% of the TRRs)	
	iviajor ivictabolites (>1076 of the	TKKS) I	viiioi ivictabo	intes (1070 of the TRRs)	
[phenyl-14C]	PI	- 1	AE C656948-b	an amida	
Intermediate plant	Fluopyram	£	AE C656948-7 AE C656948-8 glucoside conju nydroxy	7-hydroxy B-hydroxy ugate of AE C656948-7-	
Mature peppers (1x)	Fluopyram	1	AE C656948-b AE C656948-7 glucoside conju hydroxy		
Rest of plant	Fluopyram	1	AE C656948-7	7-hydroxy	
P	AE C656948-benzamide	1	AE C656948-8-hydroxy		
	(10.1% of the TRRs; 0.36 ppm)	8		malonic acid conjugates of AE	
[pyridyl-14C]					
Intermediate plant	Fluopyram	8 1	hydroxy	7-hydroxy 3-hydroxy ugate of AE C656948-7- onjugate of AE C656948-	
Mature peppers (4x)	Fluopyram AE C656948-PCA (19.5% of the TRRs; 0.029 ppm) AE C656948-PAA-glycoside [12.5% (isomer 1) and 19.7% (is of the TRRs; 0.019 and 0.029 pp	omer 2)	AE C656948-F AE C656948-7	PAA	
Mature peppers (1x)			None 2)		
Rest of plant	Fluopyram	1	hydroxy	ugate of AE C656948-7- onjugate of AE C656948-	

The main reactions of fluopyram metabolism in the beans are:

Hydroxylation of fluopyram leading to AE C656948-7-hydroxy and AE C656948-8-hydroxy

Conjugation of AE C656948-7-hydroxy with glucose and malonic acid

Cleavage of hydroxylated active substance leading to AE C656948-benzamide and AE C656948-carboxylic acid

Supplemental Cell Culture Study – Apple

PMRA# 1599640

The metabolism of fluopyram was investigated in heterotrophic plant cell suspension cultures from apple fruit following incubation with [phenyl-UL-14C] and [pyridyl-2,6-14C] fluopyram, to facilitate metabolite identification and to produce radiolabeled reference compounds for the identification of metabolites in metabolism studies. Nine metabolites (AE C656948-deschloro-3-OH-glc; AE C656948-7-hydroxy; AE C656948-7-hydroxy-glc; AE C656948-hydroxy-glc; AE C656948-benzamide; AE C656948-pyridyl-hydroxyethyl; AE C656948-pyridyl-hydroxymethyl; and AE C656948-pyridyl-carboxylic acid (PCA)) were isolated and identified, which served as reference compounds for the plant and animal metabolism studies.

Proposed Metabolism in Plants

Metabolism studies conducted in four diverse crops (pepper, grapes, beans and potato) showed similar metabolic profiles, with fluopyram as a major compound in all crops. In pepper (pyridyl), potato tuber (pyridyl) and beans (both labels), where fluopyram was not the residue present at the highest level, the TRR levels of the predominant metabolites were relatively low. In pepper, TRRs of the predominant residues were 0.01-0.026 ppm; in potato tuber, 0.003-0.006 ppm and in beans, 0.008-0.077 ppm.

The metabolism in all plants was very similar. The main reactions involved were: hydroxylation of fluopyram to AE C656948-7-hydroxy and AE C656948-8-hydroxy, conjugation of hydroxylated fluopyram mainly with sugars, cleavage of the molecule leading to AE C656948-benzamide, AE C656948-pyridyl-acetic acid (PAA) and AE C656948-carboxylic acid (PCA).

The main metabolic reactions were also observed in rats. It was concluded that the plant metabolites AE C656948-7and 8-hydroxy, AE C656948-benzamide and PAA are toxicologically covered by the data of the rat studies. Tox data for label-specific metabolite PCA were provided and showed that the metabolite is of no toxicological concern.

The metabolism of fluopyram in plants is adequately documented. The residue definition for enforcement purposes in plant commodities is fluopyram. The residue definition for risk assessment purposes is fluopyram + fluopyram-benzamide in oilseeds and legumes, and fluopyram in all other crops.

Figure 1. Proposed metabolic pathways of [phenyl-UL-14C] fluopyram in grapes.

Figure 2. Proposed metabolic pathways of [pyridyl-2,6-14C] fluopyram in grapes.

Figure 3. Proposed metabolic pathways of [phenyl-UL-14C] fluopyram in potatoes.

Figure 4. Proposed metabolic pathways of [pyridyl-2,6-14C] fluopyram in potatoes.

Figure 5. Proposed metabolic pathways of [phenyl-UL-14C] fluopyram in beans.

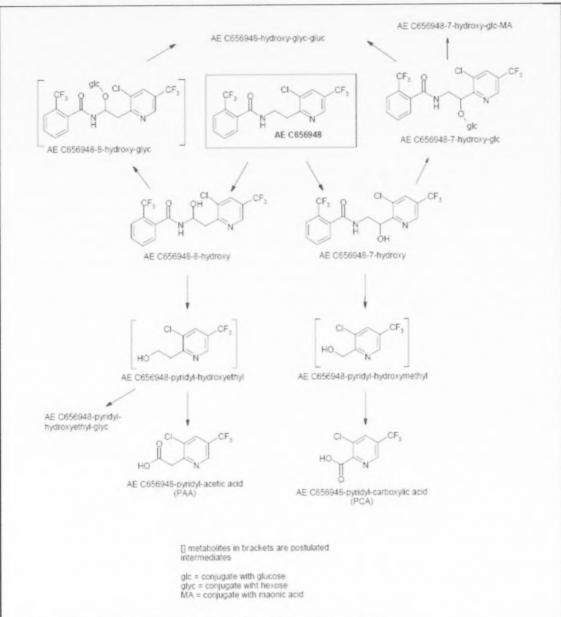


Figure 6. Proposed metabolic pathways of [pyridyl-2,6-14C] fluopyram in beans.

Figure 7. Proposed metabolic pathways of [phenyl-UL-14C] fluopyram in red bell peppers.

Figure 8. Proposed metabolic pathways of [pyridyl-2,6-14C] fluopyram in red bell peppers.

Table 18b Nature of the Residues in Plant Matrices: Confined Accumulation in Rotational Crops

Radiolahel Po	osition	[phenyl-UL-14C] fluopyram and [pyr	idyl-2,6-14C] fluopyram						
Test site		Plants were grown in vegetation halls (until cultivation of the 1st rotation) and in							
Formulation used		greenhouses (2nd and 3rd rotations)							
Formulation 1	ised	Soluble concentrate (SC) formulation	IAE C656948 formulated as SC500]						
		Soil was treated at 534 g a.i./ha (phenyl) or 514-g a.i./ha (pyridyl) and aged for 30, 139							
	ate and	and 280 days							
	dentified	and been any							
		Major Metabolites (>10% TRR)	Minor Metabolites (<10% TRR)						
		major metabolites (= 10 / v 11cts)							
		Fluopyram	AE C656948-benzoic acid						
Formulation use Application rate iming Metabolites Ide Matrix P Phenyl-UL-C14 Wheat forage 3 Wheat hay 3	150	ruopytani	AE C656948-benzamide						
			AE C656948-7-hydroxy-glc-MA (isomer 1)						
			AE C656948-8-hydroxy-glc-MA						
			AE C656948-7-hydroxy-glc						
			AE C656948-7-hydroxy						
			AE C656948-8-hydroxy						
	120	Elvenimm	AE C656948-benzoic acid						
	139	Fluopyram	AE C656948-benzamide						
			AE C656948-7-hydroxy-glc-MA (isomer 1&2)						
			AE C656948-8-hydroxy-glc-MA						
			AE C656948-7-hydroxy-glc						
			AE C656948-7-hydroxy						
			AE C656948-8-hydroxy						
	200	C1	AE C656948-benzamide						
	280	Fluopyram	AE C656948-7-hydroxy-glc-MA (isomer 1&2)						
2			AE C656948-8-hydroxy-glc-MA						
			AE C656948-7-hydroxy-glc						
			AE C656948-7-hydroxy						
			AE C656948-8-hydroxy						
	20	El	AE C656948-benzoic acid						
Wheat hay	30	Fluopyram	AE C656948-benzamide						
			AE C656948-7-hydroxy-glc-MA (isomer 1&2)						
			AE C656948-8-hydroxy-glc-MA						
Radiolabel Pos Test site Formulation us Application rat timing Metabolites Id Matrix Phenyl-UL-C1 Wheat forage			AE C656948-7-hydroxy-glc						
			AE C656948-7-hydroxy						
			AE C656948-8-hydroxy						
	120	P1	AE C656948-benzoic acid						
	139	Fluopyram	AE C656948-benzamide						
			AE C656948-7-hydroxy-glc-MA (isomer 1&2)						
			AE C656948-8-hydroxy-glc-MA						
Wheat hay			AE C656948-7-hydroxy-glc						
			AE C656948-7-hydroxy						
			AE C656948-8-hydroxy						
Wheat hay 3	200	Ch. course	AE C656948-benzamide						
	280	Fluopyram	AE C656948-7-hydroxy-glc-MA (isomer 1&2)						
		AE C656948-7-hydroxy	AE C656948-8-hydroxy-glc-MA						
			AE C656948-7-hydroxy-glc						
			AE C656948-8-hydroxy						

Wheat straw	30	Fluopyram	AE C656948-benzamide
W Hell Straw	100	, 100p; 1111	AE C656948-7-hydroxy-glc-MA (isomer 1&2)
			AE C656948-8-hydroxy-glc-MA
			AE C656948-7-hydroxy-glc
			AE C656948-7-hydroxy
			AE C656948-8-hydroxy
	139	Fluorumm	AE C656948-benzamide
	139	Fluopyram	AE C656948-7-hydroxy-glc-MA (isomer 1&2)
			AE C656948-8-hydroxy-glc-MA
			AE C656948-7-hydroxy-glc
		1	AE C656948-7-hydroxy
			AE C656948-8-hydroxy
	280	Fluopyram	AE C656948-benzamide
		AE C656948-7-hydroxy	AE C656948-7-hydroxy-glc-MA (isomer 1&2)
			AE C656948-7-hydroxy-glc
			AE C656948-8-hydroxy
Wheat grain	30	Fluopyram	AE C656948-benzoic acid
0			AE C656948-benzamide
			AE C656948-7-hydroxy-glc-MA (isomer 1)
			AE C656948-7-hydroxy-glc
			AE C656948-7-hydroxy
			AE C656948-8-hydroxy
	139	Fluopyram	AE C656948-benzamide
	137	AE C656948-benzoic acid	AE C656948-7-hydroxy
wise chard	280	Fluopyram	AE C656948-benzamide
	200	AE C656948-benzoic acid	AE C656948-7-hydroxy
c · 1 1	20		AE C656948-7-hydroxy-glc-MA (isomer 1)
Swiss chard	30	Fluopyram	AE C656948-7-OH-SA
		AE C656948-benzamide	AE C656948-7-bydroxy-glc
		AE C656948-7-hydroxy	AE C656948-8-hydroxy
	139	Fluopyram	AE C656948-benzamide
		AE C656948-7-OH-SA	AE C656948-7-hydroxy-glc
		AE C656948-7-hydroxy	AE C656948-8-hydroxy
	280	Fluopyram	AE C656948-7-hydroxy-glc
		AE C656948-benzamide	
		AE C656948-7-OH-SA	
		AE C656948-7-hydroxy	
Turnip tops	30	Fluopyram	AE C656948-benzoic acid
(leaves)		AE C656948-phenol-glc	AE C656948-benzamide
(AE C656948-7-hydroxy-glc-MA (isomer 1&2)
			AE C656948-7-OH-SA
			AE C656948-7-hydroxy-glc
			AE C656948-7-hydroxy
			AE C656948-8-hydroxy
	139	Fluopyram	AE C656948-benzamide
	137	AE C656948-phenol-glc	AE C656948-7-hydroxy-glc-MA (isomer 1&2)
		The Cost to phonor give	AE C656948-7-OH-SA
			AE C656948-7-hydroxy-glc
			AE C656948-7-hydroxy
			AE C656948-8-hydroxy
	200		AE C656948-7-hydroxy-glc-MA (isomer 1&2)
Wheat grain Swiss chard Turnip tops (leaves)	280	Fluopyram	
		AE C656948-benzamide	AE C656948-7-hydroxy
	1	AE C656948-phenol-glc	AE C656948-8-hydroxy

Turnip roots	30	Fluopyram	AE C656948-benzoic acid
rump roots		1	AE C656948-benzamide
			AE C656948-7-hydroxy
			AE C656948-8-hydroxy
	139	Fluopyram	AE C656948-benzamide
	139	поруган	AE C656948-7-hydroxy
			AE C656948-8-hydroxy
	280	Not extracted due to low residues (<0.0	
Duridul 2.6.C		Not extracted due to low residues (40.0	,1 mg/kg)
		Fluopyram	AE C656948-7-hydroxy-glc-MA (isomer 1)
wheat iotage	30	AE C656948-pyridyl carboxylic acid	AE C656948-7-hydroxy-glc
		AE Co30340-pyridyr carboxyric acid	AE C656948-7-hydroxy
Pyridyl-2,6-C1 Wheat forage	139	Fluorymm	AE C656948-methyl-sulfoxide
	139	Fluopyram	AE C656948-pyridyl carboxylic acid
			AE C656948-7-hydroxy-glc-MA (isomer 1&2)
			AE C656948-8-hydroxy-glc-MA
			AE C656948-7-hydroxy-glc
			AE C656948-7-hydroxy
			AE C656948-8-hydroxy
	280	Fluopyram	AE C656948-methyl-sulfoxide
			AE C656948-pyridyl carboxylic acid
			AE C656948-7-hydroxy-glc-MA (isomer 1&2)
			AE C656948-8-hydroxy-glc-MA
			AE C656948-7-hydroxy-glc
			AE C656948-7-hydroxy
Wheat hay	30	Fluopyram	AE C656948- pyridyl carboxylic acid
			AE C656948-7-hydroxy-glc-MA (isomer 1&2)
			AE C656948-8-hydroxy-glc-MA
			AE C656948-7-hydroxy-glc
			AE C656948-7-hydroxy
			AE C656948-8-hydroxy
	139	Fluopyram	AE C656948-methyl-sulfoxide
	137		AE C656948-pyridyl carboxylic acid
Wheat hay 3			AE C656948-7-hydroxy-glc-MA (isomer 1&2)
			AE C656948-8-hydroxy-glc-MA
			AE C656948-7-hydroxy-glc
			AE C656948-7-hydroxy
			AE C656948-8-hydroxy
	280	Chromison	AE C656948-methyl-sulfoxide
	280	Fluopyram	AE C656948-pyridyl carboxylic acid
			AE C656948-7-hydroxy-glc-MA (isomer 1&2)
			AE C656948-8-hydroxy-glc-MA
			AE C656948-7-hydroxy-glc
			AE C656948-7-hydroxy
Wheat straw	30	Fluopyram	AE C656948-pyridyl carboxylic acid
			AE C656948-7-hydroxy-glc-MA (isomer 1&2)
			AE C656948-8-hydroxy-glc-MA
			AE C656948-7-hydroxy-glc
			AE C656948-7-hydroxy
Vheat straw 3			AE C656948-8-hydroxy
	139	Fluopyram	AE C656948-methyl-sulfoxide
			AE C656948-7-hydroxy-glc-MA (isomer 1&2)
Wheat straw			AE C656948-8-hydroxy-glc-MA
			AE C656948-7-hydroxy-glc
			AE C656948-7-hydroxy
			AE C656948-8-hydroxy

	280	Fluopyram AE C656948-7-hydroxy	AE C656948-methyl-sulfoxide AE C656948-7-hydroxy-glc-MA (isomer 1&2)		
			AE C656948-8-hydroxy-glc-MA AE C656948-7-hydroxy-glc		
			AE C656948-8-hydroxy		
Wheat grain	30	Fluopyram AE C656948- pyridyl carboxylic acid	AE C656948-methyl-sulfoxide AE C656948-7-hydroxy		
	139	Fluopyram AE C656948-methyl-sulfoxide AE C656948-pyridyl carboxylic acid	AE C656948-7-hydroxy		
	280	Fluopyram AE C656948-methyl-sulfoxide AE C656948-pyridyl carboxylic acid	AE C656948-7-hydroxy		
Swiss chard	30	Fluopyram AE C656948-7-hydroxy	AE C656948-methyl-sulfoxide AE C656948-pyridyl carboxylic acid AE C656948-7-OH-SA AE C656948-7-hydroxy-glc AE C656948-8-hydroxy		
	139	Fluopyram AE C656948-7-OH-SA AE C656948-7-hydroxy	AE C656948-methyl-sulfoxide AE C656948-pyridyl carboxylic acid AE C656948-7-hydroxy-glc AE C656948-8-hydroxy		
	280	Fluopyram AE C656948-7-OH-SA AE C656948-7-hydroxy	AE C656948-methyl-sulfoxide AE C656948-pyridyl carboxylic acid AE C656948-7-hydroxy-glc AE C656948-8-hydroxy		
Turnip tops (leaves)	AE C656948- pyridyl carboxylic acid Fluopyram AE C656948-methyl-sulfoxide AE C656948-methyl-sulfoxide AE C656948-methyl-sulfoxide AE C656948-methyl-sulfoxide AE C656948-methyl-sulfoxide AE C656948-methyl-sulfoxide AE C656948-pyridyl carboxylic acid Fluopyram AE C656948-7-hydroxy Fluopyram AE C656948-7-OH-SA AE C656948-7-hydroxy Fluopyram AE C656948-7-hydroxy Important AE C656948-7-hydroxy Fluopyram AE C656948-phenol-glc Fluopyram AE C656948-phenol-glc Fluopyram AE C656948-phenol-glc	AE C656948-pyridyl carboxylic acid AE C656948-7-hydroxy-glc-MA (isomer 1&2) AE C656948-7-OH-SA AE C656948-7-hydroxy-glc AE C656948-7-hydroxy AE C656948-8-hydroxy			
	139		AE C656948-pyridyl carboxylic acid AE C656948-7-hydroxy-glc-MA (isomer 1&2) AE C656948-7-OH-SA AE C656948-7-hydroxy-glc AE C656948-7-hydroxy AE C656948-8-hydroxy		
	280		AE C656948-pyridyl carboxylic acid AE C656948-7-hydroxy-glc-MA (isomer 1&2) AE C656948-7-OH-SA AE C656948-7-hydroxy-glc AE C656948-7-hydroxy AE C656948-8-hydroxy		
Turnip roots	30	Fluopyram	AE C656948-pyridyl carboxylic acid AE C656948-7-hydroxy AE C656948-8-hydroxy		
	139	Fluopyram	AE C656948-7-hydroxy		
			AE C656948-7-hydroxy		

Proposed Metabolism in Rotational Crops

Confined rotational crop studies conducted at \sim 500 g a.i./ha on three diverse crops (wheat, Swiss chard and turnips) showed similar metabolic profiles as the ones observed in primary crops (pepper, grapes, beans and potato), with fluopyram being a major compound in all crops over all plant-back intervals. The TRRs for wheat grain, Swiss chard and turnip roots were 0.57 ppm or less and declined with the increase of PBIs. Except for fluopyram, the highest single compound measured in wheat grain, Swiss chard and turnip roots amounted to 0.23 ppm or less.

Based on the results of the confined rotational crops studies, the following predominant metabolites were observed: Wheat grain: Fluopyram, fluopyram-PCA and fluopyram-methyl sulfoxide

Swiss chard: Fluopyram, fluopyram-7-hydroxy, fluopyram-7-OH-SA (sulphate conjugate) and fluopyram-benzamide

Turnip root: Fluopyram

The residue definition for enforcement purposes is fluopyram in rotational crops.

The residue definition for risk assessment purposes is fluopyram + fluopyram-benzamide in rotational oilseeds and legumes, and fluopyram in all other rotational crops.

Figure 9. Proposed metabolic pathways of [phenyl-UL-14C] fluopyram in rotational crops.

Figure 10. Proposed metabolic pathways of [pyridyl-2,6-14C] fluopyram in rotational crops.

Table 18c Nature of the Residues in Livestock

Nature of the Residue in Laying Hen

PMRA # 1599784 and 1599792

Six laying hens (White Leghorn) were administered a single daily oral dose (in the morning by gavage using a syringe) for 14 consecutive days with either 2.03 mg per kg body weight per day (corresponding to 26.42 mg a.i./kg feed/day) for [phenyl-UL-¹⁴C]fluopyram or 2.02 mg per kg body weight per day (corresponding to 25.96 mg a.i./kg feed/day) for [pyridyl-2,6-¹⁴C]fluopyram. Animals were sacrificed about 24h after the last dose.

Phenyl Radiolabel: The overall recovery (sum of radioactivity in the excreta, eggs as well as tissues) was 94.83% of the total administered dose. The majority of the radioactivity (82.67% of the total dose) was detected in the excreta collected before sacrifice. An amount of 4.34% of the total dose was detected in the eggs. At sacrifice the compound-related residues in the edible organs and tissues amounted to 7.83% of the total dose.

The most important metabolic reaction in the laying hen was the cleavage of the aliphatic chain, yielding the major metabolite AE C656948-benzamide. A second major metabolic reaction involved the hydroxylation of the aliphatic chain followed by elimination, yielding the olefines. Hydrolysis of the amide to a carboxylic acid group was observed as a minor reaction.

Pyridyl Radiolabel: The overall recovery was 95.55% of the total administered dose. The majority of the radioactivity (94.71% of the total dose) was detected in the excreta collected before sacrifice. An amount of 0.36% of the total dose was detected in the eggs. At sacrifice the compound-related residues in the edible tissues collected from the hens amounted to 0.48% of the total dose.

The metabolic reactions in the laying hen were hydroxylation of the aliphatic chain followed by elimination, as well as oxidative cleavage of the aliphatic chain.

The results are in very good agreement with the results from the laying hen metabolism study with [phenyl-UL-14C]

fluopyram. The metabolism of fluopyram in hens is well understood.

Matrices	% of Administered Dose							
	[pho	enyl-14C]	[pyric	lyl- ¹⁴ C]				
	TRRs (ppm) (mean of 6 hens)	% of Administered Dose	TRRs (ppm) (mean of 6 hens)	% of Administered Dose				
Excreta (day 1-14)	10.655	82.67	12.642	94.71				
Total Body Muscle	3.290	4.94	0.831	0.10				
Total Body Fat	1.696	0.76	0.498	0.22				
Total Body Skin	2.533			0.02				
Liver			0.538	0.05				
Kidney	5.759		0.242	0.01				
Eggs (day 1-14)	ggs (day 1-14) 2.870		0.235	0.36				
Eggs (day 1-6)	1.811		0.156					
Eggs (day 7-14)	3.581		0.286					
Total		94.83		95.55				
Metabolites identified		es (>10% of the TRRs)	Minor Metabolites (<10% of the TRRs)					
Radiolabel Position	[phenyl-14C]	[pyridyl-14C]	[phenyl-14C]	[pyridyl-14C]				
Eggs	AE C656948-benzamide	Fluopyram AE C656948-Z-olefine	Fluopyram AE C656948-Z-olefine	AE C656948-E-olefine AE C656948-PAA AE C656948-7-hydroxy				
Muscle	AE C656948-benzamide	AE C656948-Z-olefine	AE C656948-Z-olefine	Fluopyram AE C656948-E-olefine				
Fat	AE C656948-benzamide AE C656948-Z-olefine	Fluopyram AE C656948-Z/E-olefine	Fluopyram AE C656948-E-olefine	None				
Liver	AE C656948-benzamide	AE C656948-E-olefine	AE C656948-Z/E-olefine AE C656948-benzoic acid	AE C656948-Z-olefine AE C656948-PAA AE C656948-7-hydroxy				

Nature of the Residue in Lactating Goat

PMRA # 1599783 and 1599791

One lactating goat (Bunte deutsche Edelziege) was administered a single daily dose via a gelatine capsule on five consecutive days with either 1.91 mg [phenyl-UL-¹⁴C] fluopyram per kg body weight per day (corresponding to 46.26 mg a.i./kg feed/day) or with 2.0 mg [pyridyl-2,6-¹⁴C] fluopyram per kg body weight per day (corresponding to 44.62 mg a.i./kg feed/day). The animals were sacrificed at about 24h after the last dose.

Phenyl Radiolabel: The overall recovery (sum of radioactivity in the excreta, milk as well as organs and tissues) was 93.46% of the total administered dose. The high urinary excretion during the whole testing period and the findings in the tissues suggest that a considerable amount from each oral dose was bioavailable. Up to the time of sacrifice, the excretion accounted for about 88.31% of the total dose. A high portion of 52.62% was found in the urine and 35.69% in the feces.

The metabolic reactions of [phenyl-UL-14C]fluopyram detected in the lactating goat were:

- hydroxylation of the ethylene bridge of the molecule resulting in AE C656948-7-hydroxy, AE C656948-8-hydroxy, and a dihydroxylated compound,
- hydroxylation of the phenyl ring leading to AE C656948-phenol,
- · conjugation of the hydroxylated metabolites with glucuronic acid,
- climination of water from compounds hydroxylated in the ethylene bridge leading to AE C656948-Z-olefine and E-olefine, E- and Z-olefine can isomerise into each other,
- cleavage of the aliphatic chain to form AE C656948-benzamide,
- hvdroxylation of AE C656948-benzamide followed by conjugation with sulphate.

Pyridyl Radiolabel: The overall recovery was 81.89% of the total administered dose. Up to the time of sacrifice, the excretion accounted for 80.95% of the total dose. A high portion of 52.33% was found in the urine and 28.62% in the feces.

The metabolic reactions of [pyridyl-2,6-14C]fluopyram detected in the lactating goat were:

- hydroxylation of the ethylene bridge of the molecule resulting in AE C656948-7-hydroxy, AE C656948-8-hydroxy, and a dihydroxylated compound,
- hydroxylation of the phenyl ring leading to AE C656948-phenol,
- conjugation of the hydroxylated metabolites with glucuronic acid,
- elimination of water from compounds hydroxylated in the ethylene bridge leading to AE C656948-Z-olefine and E-olefine (E- and Z-olefine can isomerize into each other).
- molecular cleavage to AE C656948-pyridyl-hydroxyethyl followed by conjugation with glucuronic acid,
- oxidation of AE C656948-pyridyl-hydroxyethyl to AE C656948-pyridyl-acetic acid.

The results are in very good agreement with the results from the lactating goat metabolism study with [phenyl-UL-

Matrices	% of Administered Dose (AD)								
	[pheny	1-14C]	[pyridy	yl- ¹⁴ C]					
	TRRs (ppm)	% of AD	TRRs (ppm)	% of AD					
Urine (0-120 h)	29.717	52.62	13.682	52.33					
Feces (0-120 h)	7.258	35.69	5.444	28.62					
Total Body Muscle	0.737	2.31	0.042	0.12					
Total Body Fat	0.399	0.50	0.372	0.42					
Kidney	2.295	0.07	0.403	0.01					
Liver	8.379	1.71	1.427	0.31					
Milk (0-120 h)	0.259	0.56	0.032	0.08					
Morning milk	0.276								
Evening Milk	0.228		0.053						
Total		93.46		81.89					

Metabolites identified	Major Metabolites	(>10% of the TRRs)	Minor Metabolites (<10% of the TRRs)		
Radiolabel Position	[phenyl-14C]	[pyridyl-14C]	[phenyl-14C]	[pyridyl-14C]	
Milk	AE C656948-benzamide	Fluopyram AE C656948-Z-olefine AE C656948-7-hydroxy	Fluopyram AE C656948-Z-olefine AE C656948-7-hydroxy AE C656948-7-OH-GA AE C656948-benzamide-SA	AE C656948-7-OH-GA AE C656948-8-OH-GA AE C656948-E-olefine	
Muscle	AE C656948-benzamide	Fluopyram AE C656948-Z-olefine AE C656948-7-hydroxy	AE C656948-7-hydroxy AE C656948-7-OH-GA AE C656948-benzamide-SA	AE C656948-7-OH-GA AE C656948-8-OH-GA AE C656948-E-olefine	
Fat	Fluopyram AE C656948-benzamide AE C656948-7-olefine AE C6		AE C656948-E-olefine AE C656948-7-hydroxy	AE C656948-E-olefine	
Liver	AE C656948-benzamide	AE C656948-7-OH-GA	Fluopyram AE C656948-Z/E-olefine AE C656948-7-hydroxy AE C656948-8-OH-GA AE C656948-8-OH-GA AE C656948-benzamide-8A AE C656948-phenol-GA	Fluopyram AE C656948-phenol-GA AE C656948-di-OH-GA AE C656948-8-OH-GA AE C656948-E-olefine AE C656948-7-hydroxy	
Kidney	AE C656948-benzamide	AE C656948-7-OH-GA	Fluopyram AE C656948-7-hydroxy AE C656948-7-OH-GA AE C656948-8-OH-GA AE C656948-benzamide-SA AE C656948-phenol-GA AE C656948-di-OH-GA	AE C656948-PAA AE C656948-hydroxyethyl-GA AE C656948-phenol-GA AE C656948-di-OH-GA AE C656948-8-OH-GA AE C656948-E-olefine AE C656948-7-hydroxy	

Proposed Metabolism in Livestock

The metabolism of fluopyram in goat and hen is very similar. The main reactions involved are:

- hydroxylation of fluopyram to AE C656948-7-hydroxy and AE C656948-8-hydroxy,
- elimination of water from compounds hydroxylated in the ethylene bridge leading to AE C656948-Z/E-olefines,
- cleavage of the molecule leading to AE C656948-benzamide and AE C656948-pyridyl-acetic acid (PAA),
- conjugation of the hydroxylated fluopyram mainly with glucuronic acid.

The metabolic pathways were similar to the ones in rat, except for the 2 isomers of fluopyram-(Z/E)-olefine which were predominant metabolites in both hen and goat matrices [not seen in rat metabolism studies; seen minimally (≤0.007 ppm) in the rat organ depletion study, in liver, kidney and perirenal fat]. It was concluded that metabolites AE C656948-benzamide and PAA are toxicologically covered by the data of the rat studies. Based on similar structure to fluopyram (and fluopicolide which is less toxic than fluopyram), fluopyram-(Z/E)-olefines are considered to be not more toxic.

The metabolism of fluopyram in animals is adequately documented. The residue definition for enforcement purposes in animal commodities is fluopyram including the metabolite fluopyram-benzamide (expressed as parent equivalent). The residue definitions for risk assessment purposes are:

- In poultry tissues and eggs: Fluopyram including the metabolites fluopyram-benzamide and fluopyram-olefines (total of 2 isomers) (expressed as parent equivalent)
- In ruminant tissues and milk: Fluopyram including the metabolites fluopyram-benzamide, fluopyram-olefines
 (total of 2 isomers) and fluopyram-7-hydroxy (expressed as parent equivalent). {Fluopyram-7-hydroxy was not
 analyzed in the feeding studies; it can be considered by a ratio (conversion factor) derived from the goat
 metabolism study with fluopyram as reference to determine the input value for risk assessment.}

Figure 11. Proposed metabolic pathways of [phenyl-UL-14C] fluopyram in laying hen.

Figure 12. Proposed metabolic pathways of [pyridyl-2,6-14C] fluopyram in laying hen.

Figure 13. Proposed metabolic pathways of [phenyl-UL-14C] fluopyram in lactating goat.

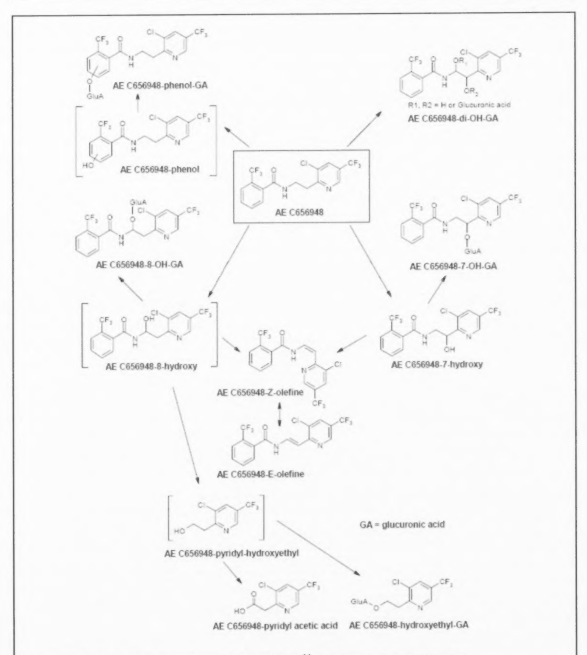


Figure 14. Proposed metabolic pathways of [pyridyl-2,6-14C] fluopyram in lactating goat.

Table 18d Freezer Storage Stability

Freezer Storage Stability PMRA# 1599821, 1784472, 1983731, 1599801, 1804905, 1983732

Residues of fluopyram and the metabolite fluopyram-benzamide are stable for up to 36 months at ≤-18°C in lettuce head, wheat grain, rape seed, dry pea seed and orange.

Stability of Other Metabolites:

Fluopyram-pyridyl-acetic acid: up to 36 months in/on lettuce head, wheat grain, rape seed and dry pea seed.

Fluopyram-pyridyl-carboxylic acid: up to 36 months in dry pea seed, rape seed and orange.

Fluopyram-7-hydroxy: up to 36 months in/on wheat grain and lettuce.

Table 18e Crop Field Trials and Residue Decline

Crop Field Trials & Residue Decline – Potatoes PMRA# 1654363

Sixteen residue trials (14 harvest and 2 decline) were conducted in 2006 (in NAFTA Growing Regions 1, 2, 3, 5, 9 and 11) on potatoes, the representative crop of Crop Group 1C. At each test location, potatoes were treated with two foliar spray applications of AE C656948 500 SC at a rate of 250 g a.i./ha/application with a 3- to 5-day application interval for a total seasonal rate of 500 g a.i./ha. The applications were made at BBCH growth stage 45 to 93 (BBCH 45: 50% of total final tuber mass reached; BBCH 93: most of leaves yellowish). Mature potato tubers were harvested at a PHI of 6-7 days. All applications were made using ground-based equipment.

At the PHI of 6-7 days, residues of fluopyram ranged from <0.01 ppm to 0.017 ppm in potato tubers (quantifiable residues were observed in only one out of the 16 trials). Fluopyram residues from both decline trials were less than the LOQ (<0.01ppm) at all time points (PHIs of 0, 3, 7, 14 and 21 days) except for the last time point of one decline trial where residues were slightly above the LOQ.

Commodity	Total Appl. Rate (g a.i./ha)	PHI	Fluopyram Residue Levels (ppm)						
•			n	Min	Max	HAFT	Median (STMdR)	Mean (STMR)	Std. Dev.
Potato tubers	500	6-7	32	< 0.01	0.017	0.016	< 0.01	< 0.01	0.004
Crop Field T	rials & Residue Dec	line – Su	gar be	ets			PMRA# 165	54364	

Twelve residue trials (11 harvest and 1 decline) were conducted in 2006 (in NAFTA Growing Regions 5, 7, 8, 9, 10 and 11) on sugar beets. At each test location, sugar beets were treated with two foliar spray applications of AE C656948 500 SC at a rate of 250 g a.i./ha/application with a 7-day application interval for a total seasonal rate of 500 g a.i./ha. The first application was made at BBCH growth stage 49 (BBCH 49: expansion complete, typical form and size of roots reached). Mature crops were harvested at a PHI of 5-7 days. All applications were made using ground-based equipment.

At the PHI of 6-7 days, residues of fluopyram ranged from 0.01 to 0.05 ppm in sugar beet roots, and from 0.27 to 18.7 ppm in sugar beet tops. In the residue decline trial, samples were harvested at PHIs of 0, 6, 13, 19 and 27 days. Mean residue level dropped from 0.07 ppm to 0.01 ppm in sugar beet roots and from 9.50 ppm to 0.04 ppm in sugar beet tops between PHIs of 0 and 27 days.

Commodity	Total Appl. Rate	PHI (days)	Fluopyram Residue Levels (ppm)						
	(g a.i./ha)		n	Min	Max	HAFT	Median (STMdR)	Mean (STMR)	Std. Dev.
Sugar beet roots	500	6-7	24	0.013	0.050	0.040	0.026	0.029	0.011
Sugar beet tops			24	0.273	18.703	16.510	0.803	3.299	4.888

Crop Field Trials & Residue Decline - Dry beans and peas

PMRA# 1661215

Nine residue trials (8 harvest and 1 decline) were conducted in 2006 (in NAFTA Growing Regions 5, 7, 8, 9, 10 and 11) on dry beans. At each test location, dry beans and peas were treated with two foliar spray applications of AE C656948 500 SC at a rate of 250 g a.i./ha per application for a total seasonal rate of 500 g a.i./ha. All applications were made using ground-based equipment.

Each trial had two treated plots (TRTD1 and TRTD2). In plot TRTD1 of the dry bean trials, the 1st application was made at a BBCH growth stage between 28 (eight side shoots detectable) and 59 (first petals visible, still closed). The 2nd application to TRTD1 was made 5-8 days later, and forage was harvested at a 0-day PHI at a target growth stage between BBCH 30 and 59. In plot TRTD2 of the dry bean trials, the 1st application was made at a BBCH growth stage between 67 (flowering declining) and 86 (60% of pods ripe and dark, seeds dry and hard). The 2nd application to TRTD2 was made 5-7 days later, and hay was harvested at a 0-day PHI at a target growth stage between BBCH 85 to 89. Seed was also harvested (plants cut from the ground) from plot TRTD2 at a 13- to 14-day PHI (except one trial with a 0-day PHI) at a target BBCH 89 growth stage. Hay and seed were allowed to dry to commercial dryness prior to sampling.

At the PHI of 13-14 days, residues of fluopyram ranged from < 0.01 to 0.08 ppm in dry beans and 0.03 to 0.35 ppm in dry peas. In the residue decline trials, seed samples were harvested at PHIs of 0, 7, 14, 17-18 and 22-24 days. Mean residue level dropped from 0.052 ppm to 0.017 ppm in dry beans between PHIs of 0 and 22 days. Residues remained approximately the same in dry peas at 0.036 ppm and 0.026 ppm between PHIs of 0 and 24 days.

The maximum fluopyram residues in dry bean forage at 0-day PHI were 25.4 ppm. The maximum fluopyram residues in dry pea vines at 0-day PHI were 11.1 ppm. The maximum fluopyram residues in dry bean hay harvested at 0-day PHI were 37.7 ppm. The maximum fluopyram residues in dry pea hay harvested at 0-day PHI were 49.4

Commodity	Total Appl. Rate	PHI	Fluopyram Residue Levels (ppm)						
	(g a.i./ha)	(days)	s) n	Min	Max	HAFT	Median (STMdR)	Mean (STMR)	Std. Dev.
Dry beans	500	13-14	18	< 0.01	0.076	0.068	0.012	0.023	0.022
Dry peas			10	0.03	0.350	0.35	0.058	0.130	0.13
	rials & Residue Dec	line - Mo	elons				PMRA# 160	51219	

Six residue trials (5 harvest and 1 decline) were conducted in 2007 (in NAFTA Growing Regions 2, 5, 6 and 10) on muskmelon. At each test location, melons were treated with two foliar spray applications of AE C656948 500 SC at a rate of 250 g a.i./ha per application with a 5- to 6-day application interval for a total seasonal rate of 500 g a.i./ha. The 1st application to melons was made at BBCH growth stage between 71 (first fruit on main stem has reached typical size and form) and 89 (fully ripe). Mature crops were harvested at a PHI of 0 day. All applications were made using ground-based equipment.

At the PHI of 0 day, residues of fluopyram ranged from 0.07 to 0.53 ppm in muskmelons. In the residue decline trials, samples were harvested at PHIs of 0, 1, 3, 7 and 10 days. Residues remained approximately the same through the 10 days in muskmelons at 0.076 ppm to 0.107 ppm. The practice of peeling muskmelon fruit treated by broadcast foliar spray reduced the total fluopyram residues in muskmelon, giving a processing factor of 0.04X.

Commodity	Total Appl. Rate	PHI (days)	Flu	Fluopyram Residue Levels (ppm)								
Commounty	(g a.i./ha)		n	Min	Max	HAFT	Median (STMdR)	Mean (STMR)	Std. Dev.			
Muskmelon	500	0	12	0.069	0.529	0.439	0.192	0.217	0.156			
Crop Field T	rials & Residue Dec	line - App	oles				PMRA# 16	70088				

Seventeen residue trials (14 harvest and 3 decline) were conducted in 2006 and 2007 (in NAFTA Growing Regions 1, 2, 5, 9, 10 and 11) on apples. At each test location, apple trees were treated with two foliar spray applications of AE C656948 500 SC at a rate of 250 g a.i./ha per application with a 5- to 7-day application interval for a total seasonal rate of 500 g a.i./ha. For all trials, there was one treated plot, which received a low volume (concentrate) spray solution, with spray volumes of 368-671 L/ha. For 12 of the apple trials, there was a second treated plot, which received a high volume (dilute) spray solution, with spray volumes of 1941-2860 L/ha. The first application was made between BBCH growth stage 78 to 89 (BBCH 78: fruit about 80% final size; BBCH 89: fruit ripe for consumption). Mature apples were harvested at a PHI of 7 days. All applications were made using ground-based equipment.

At a PHI of 7 days, residues of fluopyram ranged from 0.04 to 0.25 ppm in apples treated with a concentrated spray and from 0.06 to 0.26 ppm in apples treated with a dilute spray. In the residue decline trials, samples were harvested at PHIs of 0, 3, 7, 10 and 14 days. Mean residue level dropped from 0.16 ppm to 0.08 ppm in apples between PHIs of 0 and 14 days.

Commodity	Total Appl. Rate (g a.i./ha)	PHI (days)	Flu	Fluopyram Residue Levels (ppm)								
			n	Min	Max	HAFT	Median (STMdR)	Mean (STMR)	Std. Dev.			
Apple 500 (conc. spray)	0	34	0.054	0.796	0.751	0.192	0.225	0.151				
прри		7	34	0.040	0.247	0.242	0.109	0.120	0.063			
	500 (dilute spray)	0	24	0.070	0.545	0.437	0.159	0.176	0.092			
	Sou (and spray)	7	24	0.057	0.262	0.255	0.086	0.105	0.055			
	500	0	4	0.109	0.174	0.167	0.139	0.140	0.031			
200	5.00	7	4	0.061	0.107	0.101	0.083	0.084	0.021			
Crop Field T	rials & Residue Dec	line - Che	erries				PMRA# 160					

Six residue trials (5 harvest and 1 decline) were conducted in 2006 and 2007 (in NAFTA Growing Regions 1, 5, 10 and 11) on cherries. At each test location, cherries were treated with two foliar spray applications of AE C656948 500 SC at a rate of 250 g a.i./ha per application with a 5- to 8-day application interval for a total seasonal rate of 500 g a.i./ha. Mature crops were harvested at a PHI of 0 day. Spray volumes ranged from 371 to 624 L/ha for plots receiving concentrated sprays and from 1905 to 3350 L/ha for plots receiving diluted sprays. All applications were made using ground-based equipment.

At the PHI of 0 day, residues of fluopyram ranged from 0.07 to 0.64 in cherries treated with the concentrated spray and 0.15 to 1.2 ppm in cherries treated with the dilute spray. In the decline trials, samples were harvested at PHIs of 0, 3, 7, 10 and 14 days. Residues in cherries decreased with time. The normal household practice of washing and cooking cherries significantly reduced fluopyram residues in/on cherries. The processing factors calculated for the washed cherries and the washed and cooked cherries were 0.48X and 0.41X, respectively.

Commodity	Total Appl. Rate	PHI	Flu	Fluopyram Residue Levels (ppm)								
Commonly	(g a.i./ha)	(days)	n	Min	Max	HAFT	Median (STMdR)	Mean (STMR)	Std. Dev.			
Cherries	500 (conc. spray)	0	12	0.066	0.641	0.639	0.505	0.425	0.223			
Chemics	500 (dilute spray)	0	12	0.147	1.229	1.174	0.396	0.516	0.349			
Crop Field T	rials & Residue Dec	line - Gra	pes				PMRA# 159	99586				

Sixteen residue trials (15 harvest and 1 decline) were conducted in 2006 and 2007 (in NAFTA Growing Regions 1, 5, 10 and 11) on grapes. At each test location, two foliar spray applications of AE C656948 500 SC at a rate of 250 g a.i./ha (0.223 lb. a.i./A) were made to grapes at growth stages from fruit ripe for picking to fruit ripe for consumption (BBCH 87 to 89) with a 12 to 14 day application interval for a seasonal rate of 500 g a.i./ha, 3-day and 7-day PHIs on the harvest trials, and 0, 3, 7, 10 and 14 days PHIs for the decline trial.

Residues of fluopyram on grapes ranged from 0.068 ppm to 0.987 ppm at a PHI of 3 days and from 0.096 ppm to 0.950 ppm at a PHI of 7 days. The mean values were 0.458 ppm and 0.401 ppm on day 3 and day 7, respectively. For the decline trial, mean residue level dropped from 0.872 ppm at 0-day PHI to 0.672 ppm at 14-day PHI.

Commodity Total Appl. (g a.i./ha)	Total Appl. Rate	te PHI (days)	Fluc	Fluopyram Residue Levels (ppm)									
	1		n	Min	Max	HAFT	Median (STMdR)	Mean (STMR)	Std. Dev.				
Grapes	439 - 513	6-7	32	0.096	0.950	0.948	0.372	0.401	0.229				
Crop Field T	rials & Residue Dec	line – Sti	rawbe	rries			PMRA# 159	99587					

Ten residue trials (9 harvest and 1 decline) were conducted in 2007 (in NAFTA Growing Regions 1, 2, 3, 5, 10 and 12) on strawberries.

At each test location for the spray treated plot, two foliar spray applications of AE C656948 500 SC at a rate of 250 g a.i./ha were made to strawberry plants at BBCH growth stage 81 to 91 (beginning of ripening to beginning of auxiliary bud formation) with a 5-day application interval for a seasonal rate of 500 g a.i./ha and a 0-day PHI. A second treated plot received two drip irrigation applications of AE C656948 500 SC at a rate of 250 g a.i./ha with a 5-day interval and target PHIs of 0 and 7 days. In the decline trial, duplicate composite samples of strawberries were collected at 0, 3, 7, 10 and 14 days PHI following the final application, for plots treated by foliar spray application and drip line irrigation application.

For the broadcast application trials, fluopyram residues on strawberry fruit at the PHI of 0-day ranged between 0.18 ppm to 1.06 ppm. Data from the decline trial showed that residue levels in/on fruits dropped by about 49% over 14 days. For the drip irrigation application trials, fluopyram residues on strawberry fruit at the PHI of 0 day ranged from <LOQ to 0.11 ppm, and the residues at the PHI of 7 days ranged from <LOQ to 0.24 ppm. In the decline trial, the residue levels on day 0 were <0.01 ppm, and increased to about 0.03 ppm at day 10 and day 14 after last treatment. The residue levels in the drip irrigation trials were 5 to 10 times lower than those observed in the

Commodity	Total Appl. Rate (g a.i./ha)	PHI (days)	Flu	Fluopyram Residue Levels (ppm)								
			n	Min	Max	HAFT	Median (STMdR)	Mean (STMR)	Std. Dev.			
Strawberries 495 – 525 (d	495 – 525 (drip	0	20	< 0.01	0.112	0.10	0.01	0.026	0.028			
Ditanoemo	irrigation)	7	20	< 0.01	0.244	0.23	0.02	0.050	0.069			
4 b	491 – 519 (direct broadcast)	0	20	0.183	1.062	1.01	0.395	0.513	0.279			
	500 (European greenhouse)	1	8	0.12	0.79	0.79	0.27	0.35	0.26			
Crop Field Trials & Residue Decline – Tree Nuts							PMRA# 1661238					

Ten residue trials were conducted in 2006 on tree nuts. Five trials (4 harvest and 1 decline) were conducted on almonds (in NAFTA Growing Region 10) and five trials (4 harvest and 1 decline) were conducted on pecans (in NAFTA Growing Regions 2, 4, 6 and 8). At each test location, nuts were treated with two foliar spray applications of AE C656948 500 SC at a rate of 250 g a.i./ha per application with a 6- to 7-day application interval in almond and 13- to 14-day interval in pecan for a total seasonal rate of 500 g a.i./ha. Each trial had two treated plots, one for dilute spray applications and one for concentrated spray applications. Samples of mature nuts were harvested at a 14-day PHI. One trial for each of the representative crop was a decline trial where samples were harvested at PHIs of 0, 7, 14, 21 and 28 days. All applications were made using ground-based equipment.

At the PHI of 14 days, residues of fluopyram ranged from <0.01 ppm to 0.019 ppm in almond nutmeat, 1.22 ppm to 6.12 ppm in almond hulls and <0.01 ppm to 0.045 ppm in pecan. For the residue decline trials, mean residue level increase from <0.01 ppm on day 0 to 0.018 ppm on day 14 and decrease to 0.013 ppm on day 28 in almond nutmeat. For the residue decline trial in pecans, mean residue level dropped from 0.045 ppm to <0.01 ppm between PHIs of 0 and 14 days and remained <0.01 ppm through 28 days.

Commodity	Total Appl. Rate	PHI	Flu	Fluopyram Residue Levels (ppm)								
	(g a.i./ha)	(days)	n	Min	Max	HAFT	Median (STMdR)	Mean (STMR)	Std. Dev.			
Almond	500 (conc. spray)	14	10	< 0.01	0.016	0.015	< 0.01	< 0.01	< 0.01			
Nutmeat	500 (dilute spray)	14	10	< 0.01	0.019	0.018	< 0.01	< 0.01	< 0.01			
Almond	500 (conc. spray)	14	10	1.22	6.12	5.43	2.44	2.97	1.57			
Hulls	500 (dilute spray)	14	10	1.93	4.45	4.25	3.26	3.18	1.09			
Pecans	500 (conc. spray)	14	10	< 0.01	0.045	0.031	< 0.01	< 0.01	0.014			
	500 (dilute spray)	14	10	< 0.01	0.021	0.018	< 0.01	< 0.01	< 0.01			
Cron Field T	rials & Residue Dec	line – Pea	nuts				PMRA# 1661252					

Twelve residue trials (11 harvest and 1 decline) were conducted in 2007 and 2008 (in NAFTA Growing Regions 2, 3, 6 and 8) on peanuts.

At each test location, peanuts were treated with two foliar spray applications of AE C656948 500 SC at a rate of 250 g a.i./ha per application with for a total seasonal rate of 500 g a.i./ha. The interval between applications was 12 to 14 days. Applications were timed so that sampling would occur at growth stages from BBCH 89 (fully mature, nearly all pods developed to final size are ripe) to BBCH 97 (above ground parts of plant are dead). In the harvest trials, the representative commodities of peanut nutmeat and peanut hay were harvested at PHIs of 7 (-1) days. In the decline trial, samples of peanuts were collected at PHIs of 0, 3, 7, 10 and 14 days following the application. All applications were made using ground-based equipment.

At the PHI of 7 days, residues of fluopyram ranged from <0.01 ppm to 0.018 ppm in peanut nutmeat and 1.08 to 21.9 ppm in peanut hay. The decline of fluopyram residues with time in peanut nutmeat could not be assessed due to the low levels observed. Fluopyram residues declined with time in peanut hay.

Commodity Total Appl. R: (g a.i./ha)	Total Appl. Rate (g a.i./ha)	PHI (days)	Fluo	Fluopyram Residue Levels (ppm)									
			n	Min	Max	HAFT	Median (STMdR)	Mean (STMR)	Std. Dev.				
Peanut nutmeat	500	7	24	< 0.01	0.018	0.017	<0.01	<0.01	<0.01				
Peanut hay		7	24	1.078	21.88	20.66	6.19	8.72	6.78				
Crop Field Trials & Residue Decline – Bananas							PMRA# 160	61260					

Fourteen residue trials (12 harvest and 2 decline) were conducted in 2007 in Latin America on bananas. At each test location, bananas were treated with six foliar spray applications of AE C656948 500 SC at a rate of 100 g a.i./ha/application with a 5 to 11-day application interval for a total seasonal rate of 600 g a.i./ha. The first application was made at BBCH growth stage between 70 (first fruit visible) and 75 (fruits are 50% of final size). At each trial, single control and duplicate treated samples of bananas (bagged and unbagged) were harvested at commercial maturity, at a PHI of 0 day. In two trials, additional samples were collected at 0, 2-3, 5 and 6-7 day PHIs to monitor residue decline. All applications were made using ground-based equipment.

Residues of fluopyram on bananas (bagged; PHI of 0 day) ranged from <0.01 to 0.04 ppm (mean = 0.018 ppm). Residues of fluopyram on bananas (unbagged; PHI of 0 day) ranged from 0.018 to 0.526 ppm (mean = 0.164 ppm). In the residue decline trials, mean residues in unbagged bananas decreased from 0.04 ppm and 0.17 ppm at the 0-day PHI to <0.01 ppm and 0.13 ppm, respectively, at the 6-7 day PHI.

Commodity	Total Appl. Rate (g a.i./ha)	PHI	Flu	Fluopyram Residue Levels (ppm)									
		(days)	n	Min	Max	HAFT	Median (STMdR)	Mean (STMR)	Std. Dev.				
Bananas (unbagged)	600	0	28	0.018	0.526	0.510	0.144	0.164	0.140				

Table 18f Residue Data in Rotational Crops

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Residue Data in Rotational Crops - Limited Field Accumulation in Wheat,	PMRA# 1661301
Residue Data in Rotational Crops Elimited Lieu Lee	
Turnip and Mustard Greens	
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Three field trials each were conducted in/on rotated wheat, rotated turnips and rotated mustard greens in the US in Zones 3, 4 and 10 during the 2006 growing season.

Rotated wheat: Two foliar spray applications of AE C656948 500 SC were made to cover crops at a rate of 250 to 263 g a.i./ha/application for a total application rate of 505 to 525 g a.i./ha. The actual interval between applications was 5 to 7 days and the actual PBI ranged from 236 to 248 days. The cover crop (wheat) was harvested or destroyed within 0 to 14 days following the final application, in advance of replanting to prepare a suitable seedbed for the rotational crop.

Rotated turnips: Two foliar spray applications of AE C656948 500 SC were made to cover crops at a rate of 245 to 256 g a.i./ha/application for a total application rate of 493 to 511 g a.i./ha. The actual interval between applications was 5 to 7 days and the actual PBI ranged from 228 to 236 days. The cover crop (wheat or soybean) was harvested or destroyed within 0 to 14 days following the final application.

Rotated mustard greens: Two foliar spray applications of AE C656948 500 SC were made to cover crops at a rate of

242 to 254 g a.i./ha/application for a total application rate of 493 to 499 g a.i./ha. The actual interval between applications was 5 to 7 days and the actual PBI ranged from 228 to 236 days. The cover crop (wheat or soybean) was harvested or destroyed within 0 to 14 days following the final application.

Commodity	Total Appl. Rate	PHI (days)	Fluopyram Residue Levels (ppm)								
	(g a.i./ha)		n	Min	Max	HAFT	Median (STMdR)	Mean (STMR)	Std. Dev.		
Wheat forage			6	< 0.01	0.048	0.041	0.010	0.020	0.017		
Wheat grain	505-525		6	< 0.01	< 0.01	< 0.01	< 0.01	< 0.01	NA		
Wheat hay		8	6	0.018	0.089	0.082	0.032	0.044	0.030		
Wheat straw			6	0.011	0.12	0.12	0.031	0.056	0.054		
Turnip roots	402 511	0	6	< 0.01	< 0.01	< 0.01	< 0.01	< 0.01	NA		
Turnip tops	493-511	8	6	< 0.01	0.041	0.034	0.018	0.019	0.002		
Mustard greens	493-499	8	6	< 0.01	0.036	0.035	0.013	0.018	0.014		

Residue Data in Rotational Crops - Alfalfa

PMRA# 1654401 Twelve field trials were conducted in the US during 2007 to measure the magnitude of fluopyram residues in alfalfa

planted as a rotational crop. Two foliar spray applications of AE C656948 500 SC were made to bare soil or a cover crop (mustard) at a rate of 240 to 260 g a.i./ha/application for a total application rate of 497 to 514 g a.i./ha. The actual interval between applications was 5 to 6 days and the actual PBI ranged from 12 to 14 days. All applications were made using ground-based equipment. The fields were tilled and fertilized, or the cover crop was shredded,

disked under and the soil surface was smoothed, before seeding of alfalfa.

	Total Assal Date	PBI (days)	Flu	opyram	Residue	Levels (p	pm)		
Commodity	Total Appl. Rate (g a.i./ha)		n	Min	Max	HAFT	Median (STMdR)	Mean (STMR)	Std. Dev.
Alfalfa forage/1			24	< 0.01	0.39	0.33	0.04	0.07	0.09
Alfalfa forage/2	107.514	12-14	22	< 0.01	0.10	0.10	0.04	0.05	0.03
Alfalfa forage/3			22	0.01	0.19	0.17	0.03	0.05	0.05
Alfalfa hay/1	497-514		24	0.02	0.93	0.93	0.09	0.21	0.28
Alfalfa hay/2			22	0.01	0.36	0.35	0.11	0.13	0.11
Alfalfa hay/3			22	0.01	0.46	0.42	0.06	0.13	0.13
Residue Data in	Rotational Crops -	Cotton					PMRA# 16	61299	

Eleven field trials were conducted in the US during 2007 to measure the magnitude of fluopyram residues in cotton planted as a rotational crop. Two foliar spray applications of AE C656948 500 SC were made to bare soil at a rate of 244 to 258 g a.i./ha/application for a total application rate of 495 to 511 g a.i./ha. The actual interval between applications was 1 to 5 days and the actual PBI ranged from 12 to 14 days. All applications were made using ground-based equipment. The fields were tilled, fertilized, and rolled before planting. One trial was cancelled due to crop failure.

Commodity	Total Appl. Rate (g a.i./ha)	PBI (days)	Fluopyram Residue Levels (ppm)								
			n	Min	Max	HAFT	Median (STMdR)	Mean (STMR)	Std. Dev.		
Undelinted cottonseed	405 511	12-14	22	< 0.01	< 0.01	<0.01	0.01	0.01	NA		
Cotton gin byproducts	495-511		10	<0.01	0.02	0.02	0.01	0.01	0.01		

The following crops were originally requested as primary crops for treatment with fluopyram. It was subsequently requested that cereals, canola and soybeans be considered rotational crops only. The crop field trials were conducted according to the previously proposed Canadian GAP when treated as a primary crop.

Crop Field Trials Used as Rotational Crop Data – Field Corn and Sweet	PMRA# 1661248	
Corn		

Nineteen residue trials (four sweet corn, ten field corn and five field/sweet corn) were conducted (in NAFTA Growing Regions 1, 2, 3, 5, 6, 10, 11 and 12). At each test location, corn was treated with two foliar spray applications of AE C656948 500 SC at a rate of 250 g a.i./ha per application with a 5- to 8-day application interval for a total seasonal rate of 500 g a.i./ha. All applications were made using ground-based equipment.

Samples of corn forage and sweet corn cars (kernels plus cob with husks removed) were harvested at 0-day PHI and samples of corn stover and grain were harvested at 11- to 14-day PHIs. One trial for sweet corn and two trials for field corn were decline trials where samples were harvested at PHIs of 0-1, 3, 7, 9-10 and 13-14 days.

	Total Assal Data	DIII	Flu	opyram	Residue	Levels (ppm)			
Commodity Total Appl. Rate (g a.i./ha)	PHI (days)	n	Min	Max	HAFT	Median (STMdR)	Mean (STMR)	Std. Dev.	
Corn forage		0	32	1.56	5.52	5.15	3.29	3.52	1.14
Corn stover		11-14	30	0.70	14.69	13.40	1.69	2.61	3.09
Corn grain	500	11-14	30	< 0.01	0.020	0.018	< 0.01	< 0.01	< 0.01
Sweet corn ears		0	18	< 0.01	< 0.01	< 0.01	< 0.01	< 0.01	0
Crop Field Tria	ls Used as Rotationa	Crop Da	ita – V	Vheat ar	nd Sorgh	um	PMRA# 16	61247	

Fifteen residue trials on wheat and twelve residue trials on sorghum were conducted (in NAFTA Growing Regions 2, 4, 5, 6, 7, 8 and 11). At each test location, wheat and sorghum were treated with two foliar spray applications of AE C656948 500 SC at a rate of 250 g a.i./ha per application with a 14-day application interval for a total seasonal rate of 500 g a.i./ha. All applications were made using ground-based equipment.

Samples of wheat forage, hay, grain and straw were harvested at 12- to 15-day PHIs and samples of sorghum forage, grain and stover were harvested at 13- to 15-day PHIs. One trial each for wheat and sorghum was a decline trial where samples were harvested at PHIs of 0, 7, 14, 21 and 28 days.

	Tall A I Date	DIII	Fluopyram Residue Levels (ppm)						
Commodity Total Appl (g a.i./ha)	Total Appl. Rate (g a.i./ha)	ate PHI (days)	n	Min	Max	HAFT	Median (STMdR)	Mean (STMR)	Std. Dev.
Wheat forage	***	14	30	0.052	3.03	2.91	0.610	0.788	0.688
Wheat grain			30	0.037	0.764	0.722	0.192	0.218	0.150
Wheat hay	500		32	0.280	5.51	5.41	1.66	2.19	1.75
Wheat straw			32	0.785	12.26	11.52	4.64	4.65	3.06
Sorghum forage			24	0.18	4.10	4.08	0.858	1.13	1.12
Sorghum grain	500	14	24	0.23	3.24	3.03	0.34	0.622	0.767
Sorghum stover			24	0.19	12.15	8.63	1.01	1.65	2.49
Crop Field Trial	s Used As Rotationa	d Crop Da	ata – (Canola		PMRA#	1661254		

Eight residue trials on canola were conducted (in NAFTA Growing Regions 2, 5, 7 and 11). At each test location, canola was treated with two foliar spray applications of AE C656948 500 SC at a rate of 250 g a.i./ha per application for a total seasonal rate of 500 g a.i./ha. The interval between applications was 13 to 14 days. For all trials, the applications were made at BBCH growth stage 65 to 89 (BBCH 65: Full flowering: 50% of flowers on main raceme open; BBCH 89: Fully ripe). All applications were made using ground-based equipment.

Canola seed was harvested at a 12 to 14-day PHI at commercial maturity. One trial was a decline trial in which samples of canola seed were collected at PHIs of θ , θ , 12, 19 and 26 days following application.

	Total Annal Data	PHI (days)	Fluopyram Residue Levels (ppm)						
Commodity	Total Appl. Rate (g a.i./ha)		n	Min	Max	HAFT	Median (STMdR)	Mean (STMR)	Std. Dev.
Canola seeds	500	12-14	16	0.089	3.00	2.89	0.140	0.512	0.934
Crop Field Tri	als Used As Rotationa	d Crop D	ata –	Soybean	S	PMRA#	1661216		

Twenty residue trials on soybeans were conducted (in NAFTA Growing Regions 2, 3, 4 and 5). At each test location, soybeans were treated with two foliar spray applications of AE C656948 500 SC at a rate of 250 g a.i./ha per application for a total seasonal rate of 500 g a.i./ha. All applications were made using ground-based equipment. Each trial had two treated plots. In the treated plot from which the forage and hay were sampled, the 1st application was made at a BBCH growth stage between 14 (trifoliate leaf on the 4th node unfolded) and 75 (about 50% of pods reached final length [15–20 mm]). The 2nd application was made 5-7 days later, and forage and hay were harvested at a 6- to 7-day PHI at a target growth stage between BBCH 65 and 69. In the treated plots from which seed was sampled, the 1st application was made at a BBCH growth stage between 75 (about 50% of pods reached final length [15–20 mm]) and 88 (about 80% pods ripe, beans final color, dry and hard). The 2nd application was made 5-8 days later, with the exception of one trial that had a 14-day application interval. Seed was harvested from plot TRTDS at a 12- to 14-day PHI (except one trial with a 17-day PHI and one trial that cut the soybean plants at a 14-day PHI and sampled the seed the following day) at a target BBCH 89 growth stage. When necessary, hay was allowed to dry to commercial dryness prior to sampling. Two trials were decline trials where seed samples were harvested at PHIs of 0, 7, 21 and 28 days.

	Total Anni Data	Fluopyram Residue Levels (ppm)							
Commodity	Total Appl. Rate (g a.i./ha)	PHI (days)	n	Min	Max	HAFT	Median (STMdR)	Mean (STMR)	Std. Dev.
Soybean forage		7	40	0.320	6.19	5.70	2.53	2.62	1.48
Soybean hay	500	00	40	1.21	20.90	20.20	6.19	7.50	4.91
Soybean seed		14	40	< 0.01	0.180	0.160	< 0.01	0.021	0.036

Table 18g Residues in Processed Food and Feed

PROCESSED FOOD	AND FEED - Potato	PMRA# 1654380 (or 1661287)
Test Site	One trial in NAFTA Growing Region 5	
Treatment	Broadcast foliar applications	
Rate	Two applications at 1250 g a.i./ha for a total r	ate of 2.5 kg a.i./ha/season
End-use product	AE C656948 500 SC	
Preharvest interval	6 days	
Processed Commodity	Average Processing Factor	
Wet peel	4.3x	
Chips	0.3x	
Flakes	1.0x	
Washed tubers	0.7x	
Peeled tubers	0.2x	
Cooked tubers	0.3x	
PROCESSED FOOD	AND FEED - Sugar beet	PMRA# 1654379 (or 1661286)
Test Site	One trial in NAFTA Growing Region 5	
Treatment	Broadcast foliar applications	
Rate	Two applications at 1250 g a.i./ha for a total ra	ate of 2.5 kg a.i./ha/season
End-use product	AE C656948 500 SC	
Preharvest interval	7 days	
Processed Commodity	Average Processing Factor	
Dried pulp	1.3x	
Refined sugar	1.3x	
Molasses	0.9x	
PROCESSED FOOD A	AND FEED – Apple PMRA# 16543	83 (or 1661291), 1654393 and 1654394
North American Trials		
Test Site	One trial in NAFTA Growing Region 1	
Treatment	Broadcast foliar applications	
Rate	Two applications at 1250 g a.i./ha for a total ra	ate of 2.5 kg a.i./ha/season
End-use product	AE C656948 500 SC	
Preharvest interval	5 days	
Processed Commodity	Average Processing Factor	
Washed apples	0.7x	
Peeled apples	0.03x	
Dried apples	0.03x	
Apple juice		
	0.4x	
	0.4x 0.01x	
Applesauce		
Applesauce Wet pomace EU Trials	0.01x	
Applesauce Wet pomace EU Trials	0.01x 2.3x	and Northern Europe (Belgium and UK)
Applesauce Wet pomace EU Trials Test Site	0.01x	and Northern Europe (Belgium and UK)
Applesauce Wet pomace	0.01x 2.3x Southern Europe (Southern France and Italy) a Broadcast foliar applications	
Applesauce Wet pomace EU Trials Test Site Treatment Rate	0.01x 2.3x Southern Europe (Southern France and Italy) a	
Applesauce Wet pomace EU Trials Test Site Treatment	0.01x 2.3x Southern Europe (Southern France and Italy) a Broadcast foliar applications Four applications at 125 g a.i./ha for a total rat	
Applesauce Wet pomace EU Trials Test Site Treatment Rate End-use product	0.01x 2.3x Southern Europe (Southern France and Italy) a Broadcast foliar applications Four applications at 125 g a.i./ha for a total rat AE C656948 500 SC	

Peeled apples	0.25x			
Dried apples	0.75x			
Apple juice	0.1x			
Applesauce	0.4x			
Wet pomace	2.4x			
Dried pomace	7.7x			
PROCESSED FOOD	AND FEED - Grape PMRA# 1599645, 1599646, 1599660 and 1599584			
North American Trials				
Test Site	One trial in NAFTA Growing Region 10			
Treatment	Broadcast foliar applications			
Rate	Two applications at 1250 g a.i./ha for a total rate of 2.5 kg a.i./ha/season			
End-use product	AE C656948 500 SC			
Preharvest interval	7 days			
Processed Commodity	Average Processing Factor			
Raisins	2.4x			
Juice	0.5x			
Washed berries	0.8x			
Jelly	0.1x			
EU Trials	Vita			
Test Site	Four trials in Southern and Northern France			
Treatment	Broadcast foliar applications			
Rate	Two applications at 250 g a.i./ha for a total rate of 0.5 kg a.i./ha/scason			
End-use product	AE C656948 500 SC			
Preharvest interval	3 days			
Processed Commodity				
	Average Processing Factor			
Washed berries	0.6x 3.2x			
Wet pomace				
Dried pomace	6.4x			
Grape juice	No quantifiable residues			
Wine	0.2x			
Test Site	Southern Europe (One trial each in Spain, Portugal, Italy, Greece)			
Freatment	Broadcast foliar applications			
Rate	Two applications at 250 g a.i./ha for a total rate of 0.5 kg a.i./ha/season			
End-use product	AE C656948 500 SC			
Preharvest interval	3 days			
Processed Commodity	Average Processing Factor			
Raisins	3.7x			
	AND FEED – Strawberry PMRA# 1599587, 1599659 and 1599658			
North American Trials				
Test Site	One trial in NAFTA Growing Region 10			
Treatment	Broadcast foliar applications			
Rate	Two applications at 250 g a.i./ha for a total rate of 0.5 kg a.i./ha/season			
End-use product	AE C656948 500 SC			
Preharvest interval	0 day			
Processed Commodity	Average Processing Factor			
Washed fruit	0.7x			
Washed and cooked (~	0.7x			
am)				
EU Trials				
Γest Site	Southern and Northern France, Belgium and Spain			
Freatment	Broadcast foliar applications			
Rate	Two applications at 250 g a.i./ha for a total rate of 0.5 kg a.i./ha/season			
End-use product	AE C656948 500 SC			

Processed Commodity	Average Processing Factor	
Washed fruit	0.8x	
Preserve (incl.	0.3x	
pasteurization)	V.JA	
Jam	0.5x	
PROCESSED FOOD A		PMRA# 1654372 (or 1661275)
Test Site	One trial in NAFTA Growing Region	
Treatment	Broadcast foliar applications	
Rate	Two applications at 1250 g a.i./ha for	a total rate of 2.5 kg a i /ba/season
End-use product	AE C656948 500 SC	a total the of sie ng any me season
Preharvest interval	6 days	
Processed Commodity	Average Processing Factor	
Meal	0.2x	
Refined oil	0.3x	
Dry roasted peanuts	0.3x	
Peanut butter	0.2x	
PROCESSED FOOD A		PMRA# 1654374 (or 1661280)
Test Site	One trial in NAFTA Growing Region	
Treatment	Broadcast foliar applications	
Rate	Two applications at 1250 g a.i./ha for	a total rate of 2.5 kg a i /ha/season
End-use product	AE C656948 500 SC	a total fate of 2.3 kg a.f./fla/scasoff
Preharvest interval	14 days	
Processed Commodity	Average Processing Factor	
Bran	2.7x	
Flour	0.12x	
Middlings	0.12x 0.34x	
	0.34x 0.75x	
Shorts		
Germ	2.4x	
Aspirated grain fractions		DMD A # 4/5 4272 (4//427/)
	ND FEED - Field corn	PMRA# 1654373 (or 1661276)
Test Site	One trial in NAFTA Growing Region	3
Treatment	Broadcast foliar applications	4-4-14C2-51
Rate	Two applications at 1250 g a.i./ha for a AE C656948 500 SC	a total rate of 2.3 kg a.t./na/season
End-use product		
Preharvest interval	12 days	
Processed Commodity	Average Processing Factor	
Wet milled starch	<0.4x	
Wet-milled refined oil	0.6x	
Grits	0.5x	
Flour	0.9x	
Meal	0.8x	
Bran	2.6x	
Dry-milled refined oil	<0.4x	
Aspirated grain fractions		
PROCESSED FOOD A	ND FEED - Canola PMR	A# 1654378 (or 1661285), 1654391 and 1654395
North American Trials		
Test Site	One trial in NAFTA Growing Region	5
Treatment	Broadcast foliar applications	
Rate	Two applications at 1250 g a.i./ha for a	a total rate of 2.5 kg a.i./ha/season
End-use product	AE C656948 500 SC	
Preharvest interval	14 days	
Processed Commodity	Average Processing Factor	
Refined oil	0.01x	
Meal	0.3x	

EU Trials						
Test Site	Southern Europe (Southern France and Ita	dy) and Germany				
Treatment	Broadcast foliar applications					
Rate	Two applications at 125 g a.i./ha for a tota	ll rate of 0.25 kg a.i./ha/season				
End-use product	AE C656948 500 SC	~				
Preharvest interval	34-57 days					
Processed Commodity	Average Processing Factor					
Refined oil	1.1x					
Screwpressed oil	1.4x					
Crude oil	1.4x					
Extracted meal	0.8x					
Solvent extracted oil	1.4x					
Pomace	0.9x					
PROCESSED FOOD A	ND FEED - Soybeans	PMRA# 1654375 (or 1661282)				
Test Site	One trial in NAFTA Growing Region 5					
Treatment	Broadcast foliar applications					
Rate	Two applications at 1250 g a.i./ha for a tot	tal rate of 2.5 kg a.i./ha/season				
End-use product	AE C656948 500 SC					
Preharvest interval	13 days					
Processed Commodity	Average Processing Factor					
Meal	0.05x					
Hulls	1.3x					
Refined oil	0.02x					
Flour	0.04x					
Soy milk	0.01x					
Aspirated grain fractions	s 223x					
PROCESSED FOOD A	AND FEED - Cotton	PMRA# 1654376 (or 1661283)				
Test Site	One trial in NAFTA Growing Region 4					
Treatment	Broadcast applications to bare ground; cotton was planted with a PBI of 12 days					
Rate	Two applications at 1250 g a.i./ha for a total rate of 2.5 kg a.i./ha/season					
End-use product	AE C656948 500 SC					
Preharvest interval	Samples were collected at commercial harvest stage					
Processed Commodity	Average Processing Factor					
Meal	Desition was all OO in action and all amongs of commedities					
Hulls	Residues were <loq all="" and="" be="" commodities;="" cotton="" could="" determined.<="" factors="" in="" not="" processed="" processing="" seed="" td=""></loq>					
Refined oil	processing ractors could not be determined	d.				

Table 18h Livestock Feeding

LIVESTOCK FEEDING - Dairy cattle

PMRA# 1599761

Four treatment groups of three dairy cows each were dosed orally with fluopyram, via double-coated gelatine capsules, for 29 consecutive days at dose levels corresponding to residue intake in diet of 1 ppm, 10 ppm, 30 ppm and 100 ppm dry feed. One cow served as a control. A depuration study was conducted for the 100 ppm dosing group in which animals were sacrificed 7, 14 or 21 days after withdrawal of the dose. (One animal was excluded from the 100 ppm dose group based on reduced feed intake and no data for this animal are reported.)

Duplicate milk samples from the animals were taken before the 1st dosing (Day -7) as well as on Day 1, 2, 4, 8, 10, 13, 17, 21, 24, 26 and 29 after the 1st administered dose. Milk was additionally collected from animals in the depuration study for up to Day 50. The evening milk for each cow was frozen overnight and combined with the following morning milk sample. In addition, additional milk samples from the 100 ppm dose group were collected (Day 20/21) for processing into milk whey and milk fat (cream). Animals were sacrificed within 24 hours after the final dose, and samples of liver, muscle, kidney and fat (perirenal, subcutaneous and mesenteric) were collected for analysis.

The depuration study showed that residues of fluopyram and AE C656948-benzamide in milk and tissues decreased following the withdrawal period. Total residues of olefines in liver and kidney also decreased and total residues of olefines in muscle were below the LOQ. Total residues of olefines in subcutaneous, perirenal and mesenteric fat increased during the depuration period.

Maximum Residues of Fluopyram and Metabolites [ppm]					
Matrix	1.5 ppm dose group	14.4 ppm dose group	44.1 ppm dose group	133.1 ppm dose group	
Fluopyram					
Milk (Day 4 to	< 0.01	<0.01-0.02	0.02-0.09	0.06-0.17	
end)		(mean = 0.01)	(mean = 0.03)	(mean = 0.10)	
Skim milk	N/A	N/A	N/A	0.02	
Cream	N/A	N/A	N/A	1.4	
Fat	< 0.01	0.07	0.33	0.71	
Kidney	< 0.01	< 0.01	0.05	0.08	
Liver	0.26	0.98	2.8	4.0	
Muscle	< 0.01	< 0.01	0.04	0.03	
AE C656948-ber	nzamide				
Milk (Day 8 to	0.01-0.09	0.15-0.37	0.40-0.77	1.1-1.9	
end)	(mean = 0.02)	(mean = 0.22)	(mean = 0.54)	(mean = 1.5)	
Skim milk	N/A	N/A	N/A	1.5	
Cream	N/A	N/A	N/A	0.98	
Fat	0.01	0.33	0.45	1.1	
Kidney	0.03	0.38	0.88	1.6	
Liver	0.10	1.9	3.2	7.0	
Muscle	0.02	0.44	0.79	1.5	
Combined Residu	ues of Fluopyram and AE	C656948-benzamide			
Milk (Day 8 to	<0.02-<0.10	<0.16-0.39	0.42-0.80	1.2-2.0	
end)	(mean = 0.03)	(mean = 0.23)	(mean = 0.57)	(mean = 1.6)	
Fat	< 0.02	0.37	0.78	1.6	
Kidney	< 0.04	< 0.39	0.93	1.7	
Liver	0.36	2.3	5.3	10.9	
Muscle	< 0.03	< 0.45	0.83	1.5	
AE C656948-ole	fines				
Milk (Day 8 to	<0.02	<0.02	<0.02-0.05	0.07-0.14	
end)	<0.02	\$0.02	(mean = 0.02)	(mean = 0.10)	
Skim milk	N/A	N/A	N/A	< 0.02	
Cream	N/A	N/A	N/A	1.3	
Fat	< 0.02	0.12	0.32	0.94	
Kidney	<0.02	< 0.02	0.04	0.15	
Liver	< 0.02	0.06	0.13	0.58	
Muscle	< 0.02	< 0.02	0.03	0.04	
LIVESTOCK	FEEDING - Laying Her	n	PMRA# 1	599760	

Four treatment groups of 12 laying hens each were dosed orally with fluopyram, via feed, for 28 consecutive days at dose levels corresponding to 0.05 ppm, 0.50 ppm, 1.5 ppm and 5.0 ppm feed. Nine hens were dosed at the 0 ppm level to serve as the control group. A depuration study was conducted for the 5.0 ppm dosing group in which animals were sacrificed 8, 13 or 21 days after withdrawal of the dose.

Eggs were collected from each dose subgroup daily during the dosing period, and pooled for each subgroup per sampling day, on study days -13, -6, -1, 0, 1, 2, 5, 7, 9, 12, 14, 16, 21, 23, 26 and 28. Eggs were additionally collected from animals in the depuration study for up to Day 49. Animals were sacrificed 3-7 hours after the final dose, and samples of liver (entire organ), muscle, and overlaying skin together with any associated fat (and abdominal fat) were collected for analysis.

The depuration study showed that residues of fluopyram, AE C656948-benzamide and olefines in eggs and poultry tissues decreased following the withdrawal period.

	Maximum Residues o	f Fluopyram and Meta	bolites [ppm]	
Matrix		0.49 ppm dose group		4.8 ppm dose group
Fluopyram				
Egg (Day 21 to end)	<0.01	<0.01	<0.01	<0.01
Skin with fat	< 0.01	< 0.01	< 0.01	<0.01
Liver	< 0.01	< 0.01	< 0.01	< 0.01
Muscle	< 0.01	< 0.01	< 0.01	< 0.01
AE C656948-ben	zamide			
Egg (Day 21 to end)	<0.01	0.07-0.09 (mean = 0.08)	0.20-0.23 (mean = 0.21)	0.64-0.76 (mean = 0.71)
Skin with fat	< 0.01	0.04	0.11	0,63
Liver	0.02	0.16	0.43	1.6
Muscle	< 0.01	0.04	0.10	0.33
Combined Residu	ies of Fluopyram and AE	C656948-benzamide		
Egg (Day 21 to end)	<0.02	<0.08-<0.10 (mean = 0.09)	<0.21-<0.24 (mean = 0.22)	<0.65-<0.77 (mean = 0.72)
Skin with fat	< 0.02	< 0.05	< 0.12	< 0.64
Liver	< 0.03	< 0.17	< 0.44	<1.6
Muscle	< 0.02	< 0.05	< 0.11	< 0.34
AE C656948-ole	fines			
Egg (Day 21 to end)	<0.02	<0.02	<0.02	<0.02-0.02
Skin with fat	< 0.02	< 0.02	0.03	0.08
Liver	< 0.02	< 0.02	< 0.02	0.02
Muscle	< 0.02	< 0.02	< 0.02	0.06

Table 19 Food Residue Chemistry Overview of Metabolism Studies and Risk Assessment

PLANT S	STUDIES
RESIDUE DEFINITION FOR ENFORCEMENT Primary crops and rotational crops	Fluopyram
RESIDUE DEFINITION FOR RISK ASSESSMENT Crop Groups 6 (Legume Vegetables) and 20 (Oilseeds)	Fluopyram including the metabolite fluopyram- benzamide (expressed as parent equivalent)
All other crops	Fluopyram
METABOLIC PROFILE IN DIVERSE CROPS	Similar in grapes, potatoes, beans, red bell peppers
ANIMAL	STUDIES
RESIDUE DEFINITION FOR ENFORCEMENT Ruminants and poultry	Fluopyram including the metabolite fluopyram- benzamide (expressed as parent equivalent)
RESIDUE DEFINITION FOR RISK ASSESSMENT Poultry tissues and eggs	Fluopyram including the metabolites fluopyram- benzamide and fluopyram-olefines (total of 2 isomers) (expressed as parent equivalent)
Ruminant tissues and milk	Fluopyram including the metabolites fluopyram- benzamide, fluopyram-olefines (total of 2 isomers) and fluopyram-7-hydroxy (expressed as parent equivalent)

METABOLIC PROFILE IN ANIMALS		Similar in goat, hen, rat		
FAT SOLUBLE RESIDUE		No		
DIETARY RISK FROM FOOD	AND WATER			
	POPULATION	ESTIMATED RISK % of ACCEPTABLE DAILY INTAKE (A		
		Food Only	Food and Water	
Refined chronic dietary risk	All infants < 1 year	3.9	63.8	
	Children 1-2 years	6.3	33.4	
ADI = 0.012 mg/kg bw/day	Children 3–5 years	3.9	29.3	
Estimated about a delation	Children 6–12 years	2.0	19.5	
Estimated chronic drinking water concentration =	Youth 13-19 years	0.8	14.0	
104 μg a.i./L	Adults 20-49 years	0.8	17.8	
	Adults 50+ years	1.0	18.9	
	Females 13-49 years	0.8	17.8	
	Total population	1.3	19.6	
	POPULATION	ESTIMATED RISK % of ACUTE REFERENCE DOSE (ARfD)		
			Food and Water	
Basic acute dietary exposure	All infants < 1 year	7.0	9.1	
analysis, 95 th percentile	Children 1–2 years	8.8	9.8	
Estimated acute drinking water	Children 3–5 years	7.5	8.4	
	Children 6–12 years	5.3	5.8	
concentration = 106 µg a.i./L	Youth 13-19 years	3.5	3.9	
ARfD = 0.5 mg/kg bw	Adults 20–49 years	2.8	3,3	
	Adults 50+ years	2.2	2.8	
	Females 13–49 years	2.8	3.3	
	Total population	4.4	5,0	
Refined chronic cancer dietary risk	POPULATION	ESTIMATED RISK Adjusted for Limited 3-Year Application Pe		
		Food and Water		
Q_1 * = 0.0172 (mg/kg bw/day) ⁻¹ Estimated chronic drinking water concentration = 2.93 μ g a.i./L	Total population	1 × 10 ⁻⁶		

Table 20 Summary of Physico-Chemical Properties of Fluopyram Relevant to the Environment

Parameter	Values	Interpretation		
Water solubility (at 20°C)	pH 4 \rightarrow 15 mg/L pH 7 \rightarrow 16 mg/L pH 9 \rightarrow 15 mg/L	soluble under environmentally relevant pH conditions		
Vapour pressure/volatility	20°C → 1.2 × 10 ⁻⁶ Pa 25°C → 3.1 × 10 ⁻⁶ Pa 50°C → 2.9 × 10 ⁻⁴ Pa	non-volatile under field conditions		
Henry's Law Constant	$20^{\circ}\text{C} \rightarrow 2.98 \times 10^{-5} \text{ Pa m}^3 \text{ mol}^{-1}$	low potential for volatilization from moist soil and water surfaces		

Parameter	Values	Interpretation		
UV absorption	< 292 nm	low potential for phototransformation		
pK _a (at 23°C)	0.5	does not dissociate under environmentally relevant pH conditions		
Kow/log Kow	pH $6.5 \rightarrow 2060 / 3.3$	potential for bioaccumulation		
Stability of compound at room temperature	Stable, no decomposition			

Table 21 Fate and Behaviour in the Terrestrial Environment

Study	Label/Product	System	DT ₅₀ (days)	DT ₉₀ (days)	Kinetic Model
	1	Soils			
Hydrolysis	phenyl	stable to hydrolysis under acidic, n	eutral and alkali	ine conditions	
Soil Photolysis		stable to photolysis in soils			
Aerobic soil	phenyl	Hohenseh silt loam	221	735	SFO
		AXXa sandy loam	231	769	SFO
		Wurmwiese loam	339	>1000	SFO
		Allla loam	165	549	SFO
		Porterville sandy loam	(days) (days) (days)	>1000	SFO
		Springfield silt clay loam	654	>1000	DFOP
	pyridyl	Hohenseh silt loam	210	697	SFO
		AXXa sandy loam	464	>1000	SFO
		Wurmwiese sandy loam	250	829	SFO
		Dollendorf clay loam	162	538	SFO
		Porterville sandy loam	561	>1000	SFO
		Springfield silty clay loam	583	>1000	DFOP
Anaerobic soil	phenyl	Hoefchen silt loam	>1000	>1000	SFO
	pyridyl	Hohenseh silt loam	>1000	>1000	SFO
Field studies:	dissipation:	Burscheid, GER [silt loam]	145	>1000	DFOP
Europe		Little Shelford, UK [sandy loam]	164	>1000	DFOP
•		Staffanstorp, Sweden [loam]	386	>1000	SFO
		Vatteville, France [silt loam]	318	>1000	DFOP
		Vilobi d'Onyar, Spain [loam]	221	487	SFO
		Albaro, Italy [silt loam]			DFOP
	accumulation: fluopyram 250SC	Monheim, GER [sandy loam]	end of 1 st year:29% of 0-day 1 st application end of 2 nd year:57% of 0-day 2 nd application		
		Tarascon, France [silt loam]	end of 1st year: 53% of 0-day 1st application		
Field studies: US	accumulation: New York [loamy sand] 539 500SC North Dakota [loam] 83 Georgia [loamy sand] 24	Washington [sandy loam]	163	1.00	DFOP
		New York [loamy sand]	539	DT _{75:} >1000 DT _{90:} >1000	DFOP
		DT ₇₅ >1000 DT ₉₀ >1000	DFOP		
		Georgia [loamy sand]	24	DT _{75:} 521 DT _{90:} >1000	DFOP
		California [sandy loam]	174		DFOP

Study	Label/Product	System	DT ₅₀ (days)	DT ₉₀ (days)	Kinetic Model
Adsorption/ desorption	AE C656948	Laacherhof AXXa(sandy loam)	K _{d(ad)} : 3.80 K _{oc(ad)} : 292	K _{d(des)} : 8.27 K _{oc(des)} : 636	
		Hoefchen a. Hohenseh(silt loam)	K _{d(ad)} : 8.37 K _{oc(ad)} : 322	K _{d(des)} : 13.15 K _{oc(des)} : 506	
		Laacherhof Wurmwiese(loam)	K _{d(ad)} : 5.59 K _{oc(ad)} : 266	K _{d(des)} : 9.33 K _{oc(des)} : 444	
		Pikeville(loamy sand)	K _{d(ad)} : 3.16 K _{oc(ad)} : 288	K _{d(des)} : 6.32 K _{oc(des)} : 575	
		Stilwell(clay loam)	K _{d(ad)} : 5.06 K _{oc(ad)} : 460	K _{d(des)} : 9.17 K _{oc(des)} : 834	
	AE C656948-7- hydroxy	AIIIa(loam)	K _{d(ad)} : 1.03 K _{oc(ad)} : 94	K _{d(des)} : 3.54 K _{oc(des)} : 322	
		AXXa(sandy loam)	K _{d(ad)} :1.36 K _{oc(ad):} 91	K _{d(des)} : 3.78 K _{oc(des)} : 252	
		Hoefchen(silt loam)	K _{d(ad)} : 2.54 K _{oc(ad)} : 159	K _{d(des)} : 7.16 K _{oc(des)} : 447	
		Wurmwiese(sandy loam)	K _{d(ad)} : 1.38 K _{oc(ad)} : 86	K _{d(des)} : 3.88 K _{oc(des)} : 243	

Table 22 Fate and Behaviour in the Aquatic Environment

Study	Label	System	DT ₅₀ (days)	DT ₉₀ (days)	Kinetic model
		Aquatic systems			
Hydrolysis		stable to hydrolysis under a	cidic, neutral and a	lkaline co	onditions
Water photolysis	phenyl and pyridyl	buffer solution (pH 7)	21 and 25 52 and 63 ^a 81 and 97 ^b		SFO
	phenyl and pyridyl	natural water/sediment	21, 87 ^s and 135 ^b		SFO
Aerobic aquatic	phenyl	Angleweiher -water phase	25	280	DFOP
, , , , , , , , , , , , , , , , , , , ,		Angleweiher-total system	1190	3960	SFO
		Lawrence-water phase	14	220	DFOP
		52 and 63° 81 and 97°	3300	SFO	
	pyridyl	Angleweiher-water phase	26	290	DFOP
		Angleweiher-total system	1470	4900	SFO
		Lawrence-water phase	17	220	DFOP
		Lawrence-total system	650	2150	SFO
Anaerobic aquatic	phenyl	Lawrence-water phase	4	89	DFOP
		Lawrence-total system	1580	5240	SFO
	pyridyl	Lawrence-water phase	5	79	FOMC
		Lawrence-total system	1410	4680	SFO

^aequivalent days of sunlight in Phoenix, Arizona ^b Equivalent days of sunlight in Athens, Greece

Table 23 Maximum Concentrations of Transformation Products in Soil and Water

Property	Transformation products						
	Major	Minor					
	Soil						
Hydrolysis	None	None					
Phototransformation	None	None					
Anacrobic Biotransformation Anacrobic Biotransformation Biotransformation Field dissipation: Europe Field dissipation: US AE AE AC AC AC AC AC AC AC AC	None	AE C656948-7-hydroxy (4.2% AR)					
		AE C656948- pyridyl-carboxylic acid (0.7% AR)					
		AE C656948-methyl-sulfoxide (1.0% AR)					
iotransformation ield dissipation: Europe ield dissipation: US AE		AE C656948-benzamide (1.1% AR)					
Anaerobic Biotransformation	None	None					
Field dissipation: Europe	no	t determined					
Field dissipation: US	AE C656948-benzamide (19%)*	AE C656948-7-hydroxy (3%)**					
•	AE C656948- pyridyl-carboxylic	AE C656948-benzamide**					
	acid (16%)*	AE C656948- pyridyl-carboxylic acid**					
	Water						
Hydrolysis	None	none					
Phototransformation	pH 7 buffer: AE C656948-lactam (13% AR)	none					
	natural water/sediment system:	AE C656948-lactam (1.2% AR)					
Aerobic Biotransformation	none	none					
Anacrobic Biotransformation	none	none					

AR: applied radioactivity

Table 24 Structure and Properties of Parent Compound and Transformation Products

Common name	Chemical name (CAS)	Structure	Formula and molar mass
Fluopyram	Benzamide, N-[2-[3-chloro-5- (trifluoromethyl)-2-pyridinyl]ethyl]- 2-(trifluoromethyl)- (9CI)	CF ₃ Cl CF ₃	C ₁₆ H ₁₁ Cl F ₆ N ₂ O 396.72 g/mol
Fluopyram - 7-hydroxy	N-{2-[3-chloro-5- (trifluoromethyl)pyridin-2-yl]-2- hydroxyethyl}-2- (trifluoromethyl)benzamide	CF ₃ OH CF ₃	C ₁₆ H ₁₁ ClF ₆ N ₂ O ₂ 412.72 g/mol
Fluopyram- benzamide	2-trifluoromethyl benzamide	OF ₃ ONH ₂	C ₈ H ₆ F ₃ NO 189.15 g/mol

^{(): %} of 0-day concentration

^{*}detected only at the California site

^{**} detected at sites relevant to Canadian field use conditions

Common name	Chemical name (CAS)	Structure	Formula and molar mass
Fluopyram - pyridyl- carboxylic acid	[3-chloro-5- (trifluoromethyl)pyridin-2- carboxylic acid	HO N CF ₃	C ₇ H ₃ Cl F ₃ N O ₂ 225.26 g/mol
Fluopyram- lactame	2,9-bis(trifluoromethyl)-6,7-dihydropyrido[2,3-e][2]benzazocin-8(5H)-one	F ₃ C CF ₃	C ₁₆ H ₁₀ F ₆ N ₂ O 360.26 g/mol

Table 25 Screening Level EECs* (Luna Privilege)

Soil**	Wate	1***		
	15 cm depth	80 cm depth		
0.22 mg a.i./kg soil	0.33 mg a.i./L water	0.062 mg a.i./L water		

^{*}based on 2 application of 250 g a.i./ha each with a cumulative application rate 497.76 g

**top 30 cm soil depth and a soil bulk density of 1.5 g/cm³

Table 26 Level 1 Aquatic Eco-Scenario Modelling EECs for Fluopyram in a Water Body of 0.15 m Deep Excluding Spray Drift

Use pattern		EEC (μg a.i./L)								
	Peak	96-hour	21-day	60-day	90-day	Yearly				
2×0.25 kg a.i./ha,	at 7-day interva	ls								
Potato-PEI			261	253	252	244				

Vulnerable scenario used in this Level 1 aquatic eco-scenario modelling.

Table 27 EECs in Vegetation and Insects after a Direct Over-Spray¹ (Luna Privilege)

Matrix	EEC ^a (mg a.i./kg fw) (Max Residues)	Fresh/Dry Weight Ratios	EEC Direct Overspray (mg a.i./kg dw) (Max Residues)	EEC (Direct Overspray) (mg a.i./kg dw) (Mean Residues)
Short range grass	86.4383	3.3 ^b	285.2465	101.3026
Leaves and leafy crops	48.8730	11 ^b	537.6031	177.7200
Long grass	39.5830	4.4 ^b	174.1652	56.8704
Forage crops	48.8730	5.4 b	263.9143	87.2444
Small insects	21.0032	3.8°	79.8123	44.5107
Pods with seeds	5.2508	3.9°	20.4783	9.7666
Large insects	5.2508	3.8°	19.9532	9.5161
Grain and seeds	5.2508	3.8°	19.9532	9.5161
Fruit	5.2508	7.6 °	39.9064	19.0323

based on direct over-spray of a cumulative application rate of 403.909 g a.i./ha and a default half-life of 10 days)

^{***} cumulative application of 499.18 g a.i./ha based on a half-life of 1470 days

^a based on correlations reported by Hoerger and Kenaga (1972) and Kenaga (1973)

^b fresh to dry weight ratios from Harris (1975)

^c fresh to dry weight ratios from Spector (1956)

Table 28 Effects on Terrestrial Organisms

Organism	Exposure	xposure Test Substance End-Point Value				
		Invertebr	rates			
Earthworm (Eisenia fetida andrei)	14-d acute	AE C656948	LC ₅₀ : >1000 mg a.i./kg dw soil NOAEC: 100 mg a.i./kg dw soil EC ₅₀ : >1000 mg a.i./kg dw soil	N/A	1599606	
	14-d acute	Luna Privilege G	LC ₅₀ : >415 mg a.i/kg dw soil EC ₅₀ : >415 mg a.i./kg dw soil NOAEC: 73.87a.i./kg dw soil	N/A	1599293	
	reproduction (number of juveniles)	Luna Privilege G	EC ₅₀ : >20.3 mg a.i./kg dw soil; NOAEC: 11.4 mg a.i./kg dw soil	N/A	1599294 1599589	
Honcybees (Apis mellifera L.)	48-h acute oral	AE C656948	LC ₅₀ : >102.3 μg a.i./bee NOAEL: 102.3 μg a.i./bee LOAEL: >102.3 μg a.i./bee	relatively nontoxic	1599733	
	48-h acute contact	AE C656948	LD ₅₀ : >100 μg a.i/bee NOAEL: 100 μg a.i/bee LOAEL: >100 μg a.i./bee	relatively nontoxic		
	48-h acute contact and Oral	Luna Privilege G	Contact: LD ₅₀ : >83.2 µg a.i./bee NOAEL: 83.2 µg a.i./bee LOAEL: >83.2 µg a.i./bee Oral: LC ₅₀ : >89 µg a.i./bee NOAEL: 89 µg a.i./bee LOAEL: >89 µg a.i./bee	relatively nontoxic	1599290	
Parasitic wasp	acute	Luna Privilege G	N/A	1599729		
(Aphidius rhopalosiphi)	chronic (reproduction)		LR ₅₀ : >1008 g a.i./ha ER ₅₀ : >1008 g a.i./ha	N/A	1599291	
Predatory mite	acute	Luna Privilege G	LR ₅₀ : >1008 g a.i./ha	N/A	1599727	
(Typhlodromus pyri)	Chronic (reproduction)		NOAEL:1008 g a.i./ha	N/A	1599292	
		Other soil inve	rtebrates			
Rove beetle (Aleochara bilineata)	chronic (reproduction)	Luna Privilege G	ER ₅₀ : >1008 g a.i./ha	N/A	1599634 1599295	
Soil mite Hypoaspis aculeifer (Acari laclapidae)	14-d reproduction test	Luna Privilege G	LC ₅₀ > 415 mg a.i /kg dw soil NOEC: 415 mg a.i /kg dw soil	N/A	1599296	
Springtail Folsomia candida (Collembola isotomidae)	chronic (reproduction)	Luna Privilege G	NOEC: 103.7 mg a.i./kg dw soil	N/A	1599297	
		Birds		•		
Bobwhite quail (Colinus	acute		LD ₅₀ : >2000 mg a.i./kg bw NOAEL: <500 mg a.i./kg bw	practically non-toxic	1599536	
viriginianus)	dictary		LC ₅₀ : >4785 mg a.i./kg diet LD ₅₀ : 1845.4 mg a.i./kg bw/day LOAEC: 279 mg a.i./kg diet NOAEC: <279 mg a.i./kg diet	practically non-toxic	1599554	

Organism	Exposure	Test Substance	End-Point Value	Degree of Toxicity	PMRA#
	reproduction	AE C656948	NOAEC: 46.7 mg a.i./kg diet (survival body weight) NOAEL: 4.12 mg a.i./kg bw/day LOAEC: 75.7 mg a.i./kg diet LOAEL: 6.8 mg a.i./kg bw/day	N/A	1599605
Mallard duck (Anas platyrhynchos)	dictary	AE C656948	LC ₅₀ : >4604.5 mg a.i./kg diet LD ₅₀ : 1642.7 mg a.i./kg bw/day NOAEC: 2307.1 mg a.i./kg diet LOAEC: 4604.5 mg a.i./kg diet	practically non-toxic	1599600
	reproduction	AE C656948	NOAEC: 183 mg a.i./kg diet (survivor weights) NOAEL: 18.46 mg a.i./kg bw/day LOAEC: 428 mg a.i./kg diet	N/A	1599731
		Mamm	als		
Rat	acute oral	AE C 656948, Luna Privilege	LD ₅₀ : >2000 mg a.i./kg bw	practically non-toxic	
	dietary	AE C 656948	NOAEL: 12.5 mg a.i./kg bw/d		
	reproduction	AE C 656948	NOAEL: 13.9 mg a.i./kg bw/d		
		Vascular	plants		
Vascular plant	seedling emergence	Luna Privilege G	Monocot, most sensitive: none EC ₂₅ : > 500 g a.i./ha Dicot: most sensitive: Buckwheat (Biomass) EC ₂₅ : >500 g a.i./ha	N/A	1599302 1599591
	vegetative vigour	Luna Privilege G	Monocot, most sensitive: none EC ₂₅ : > 250 g a.i./ha Dicot: most sensitive: none EC ₂₅ : >250g a.i./ha	N/A	1599301 1599590

Table 29 Screening Level Risk Assessment to Terrestrial Organisms (Luna Privilege)

Organism	Exposure	Tox Value for RQ	EEC	RQ	
Earthworm	acute	Luna Privilege	LC50×0.5: 207.5 mg a.i./kg dw soil*	0.22 mg a.i./kg soil	0.001
(E. fetida)	reproduction	Luna Privilege	NOAEC: 11.4 mg a.i./kg dw soil	0.22 mg a.i./kg soil	0.02
Honeybees (A. mellifera L.)	acute contact	Luna Privilege	LD ₅₀ : 93.2 kg a.i./ha**	0.4039 kg a.i./ha	0.004
Predatory mite	acute	Luna Privilege	LR ₅₀ : >1008 g a.i./ha	403.9 g a.i./ha	< 0.40
(T. pyri)	chronic (reproduction)	Luna Privilege	NOAEL: 1008 g a.i./ha	403.9 g a.i./ha	0.40

^{*} with an uncertainty factor of two

Table 30 Screening Level Risk Assessment to Wild Birds

Effects	Toxicity (mg a.i./kg bw/d)	Feeding Guild (food item)	EDE (mg a.i./kg bw)	RQ
		Small Bird (0.02 kg)		
Acute	200.00*	Insectivore (small insects)	20.35	0.10
Reproduction	4.12	Insectivore (small insects)	20.35	4.94
	Med	dium Sized Bird (0.1 kg)		

^{**} LD_{50} of >83.2 μg a.i./bee converted to >93.2 kg a.i./ha (based on conversion factor of 1.12 to kg per hectare according to Atkins *et al.*(1981)

Effects Toxicity (mg a.i./kg bw/d)		I Hooding (-mild (food ifem)			
Acute	200.00*	Insectivore (small insects)	15.88	0.08	
Reproduction	4.12	Insectivore (small insects)	15.88	3.86	
	L	arge Sized Bird (1 kg)			
		Herbivore (short grass)	16.57	0.08	
Reproduction	4.12	Herbivore (short grass)	16.57	4.02	

^{*} based on an uncertainty factor of 10

Table 31 Expanded Screening Level Reproductive Risk Assessment to Wild Birds for On-Field and Off-Field Scenarios (Luna Privilege)

	Toxicity			Maximum Residues							Mean Residues							
Effects	(mg a.i./kg	Food Guild (food item)	On-	On-field		Off-field (74%)		Off-field (59%)		Off-field (6%)		ield	Off-field (74%)		Off-field (59%)		Off-field (6%)	
	bw/d)		EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ
				Sn	nall Bi	ird (0	.02 kg)										
	4.12	Insectivore (small insects)	20.35	4.94	15.06	3.66	12.01	2.91	1.22	0.30	11.35	2.75	8.40	2.04	6.70	1.63	0.68	0.17
Reproduction	4.12	Granivore (grain and seeds)	5.09	1.23	3.77	0.91	3.00	0.73	0.31	0.07	2.43	0.59	1.80	0.44	1.43	0.35	0.15	0.04
	4.12	Frugivore (fruit)	10.18	2.47	7.53	1.83	6.00	1.46	0.61	0.15	4.85	1.18	3.59	0.87	2.86	0.70	0.29	0.07
			1	Mediu	ım Siz	ed Bi	rd (0.1	kg)										
	4.12	Insectivore (small insects)	15.88	3.86	11.75	2.85	9.37	2.27	0.95	0.23	8.86	2.15	6.55	1.59	5.23	1.27	0.53	0.13
n 1 .:	4.12	Insectivore (large insects)	3.97	0.96	2.94	0.71	2.34	0.57	0.24	0.06	1.89	0.46	1.40	0.34	1.12	0.27	0.11	0.03
Reproduction	4.12	Granivore (grain and seeds)	3.97	0.96	2.94	0.71	2.34	0.57	0.24	0.06	1.89	0.46	1.40	0.34	1.12	0.27	0.11	0.03
	4.12	Frugivore (fruit)	7.94	1.93	5.88	1.43	4.69	1.14	0.48	0.12	3.79	0.92	2.80	0.68	2.23	0.54	0.23	0.06
				Lar	ge Size	ed Bir	rd (1 k	g)										
	4.12	Insectivore (small insects)	4.64	1.13	3.43	0.83	2.74	0.66	0.28	0.07	2.59	0.63	1.91	0.46	1.53	0.37	0.16	0.04
	4.12	Insectivore (large insects)	1.16	0.28	0.86	0.21	0.68	0.17	0.07	0.02	0.55	0.13	0.41	0.10	0.33	0.08	0.03	0.01
	4.12	Granivore (grain and seeds)	1.16	0.28	0.86	0.21	0.68	0.17	0.07	0.02	0.55	0.13	0.41	0.10	0.33	0.08	0.03	0.01
Reproduction	4.12	Frugivore (fruit)	2.32	0.56	1.72	0.42	1.37	0.33	0.14	0.03	1.11	0.27	0.82	0.20	0.65	0.16	0.07	0.02
	4.12	Herbivore (short grass)	16.57	4.02	12.26	2.98	9.78	2.37	0.99	0.24	5.89	1.43	4.36	1.06	3.47	0.84	0.35	0.09
	4.12	Herbivore (long grass)	10.12	2.46	7.49	1.82	5.97	1.45	0.61	0.15	3.30	0.80	2.45	0.59	1.95	0.47	0.20	0.05
	4.12	Herbivore (forage crops)	15.33	3.72	11.35	2.75	9.05	2.20	0.92	0.22	5.07	1.23	0.30	0.07	2.99	0.73	0.30	0.07

Table 32 Refined Assessment of Reproductive Risk to Wild Birds for On-Field and Off-Field Scenarios (Luna Privilege)

	Toxicity	Maximum Residues								Me	ean F	Residu	es					
Effects	(mg a.i./kg	Food Guild (food item)	On-f	ield	Off-1		Off-1	-	Off-1		On-f	ield	Off-		Off-1		Off-1	
	bw/d)		EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ
				Sı	nall B	ird (0	.02 kg)										
	6.8	Insectivore (small insects)	20.35	2.99	15.06	2.21	12.01	1.77	1.22	0.18	11.35	1.67	8.40	1.24	6.70	0.98	0.68	0.10
Reproduction	6.8	Granivore (grain and seeds)	5.09	0.75	3.77	0.55	3.00	0.44	0.31	0.04	2.43	0.36	1.80	0.26	1.43	0.21	0.15	0.02
	6.8	Frugivore (fruit)	10.18	1.50	7.53	1.11	6.00	0.88	0.61	0.09	4.85	0.71	3.59	0.53	2.86	0.42	0.29	0.04
			1	Medi	um Siz	ed Bi	rd (0.	kg)										

	Toxicity				Max	imun	Resid	lues					M	ean F	Residu	es		
Effects	(mg a.i./kg	Food Guild (food item)	On-	field	Off-		Off-1		Off-		On-	ield	Off-		Off-		Off-	
	bw/d)		EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ
	6.8	Insectivore (small insects)	15.88	2.34	11.75	1.73	9.37	1.38	0.95	0.14	8.86	1.30	6.55	0.96	5.23	0.77	0.53	0.08
	6.8	Insectivore (large insects)	3.97	0.58	2.94	0.43	2.34	0.34	0.24	0.04	1.89	0.28	1.40	0.21	1.12	0.16	0.11	0.02
Reproduction	6.8	Granivore (grain and seeds)	3.97	0.58	2.94	0.43	2.34	0.34	0.24	0.04	1.89	0.28	1.40	0.21	1.12	0.16	0.11	0.02
	6.8	Frugivore (fruit)	7.94	1.17	5.88	0.86	4.69	0.69	0.48	0.07	3.79	0.56	2.80	0.41	2.23	0.33	0.23	0.03
				Lar	ge Siz	ed Bi	rd (1 k	ig)										
	6.8	Insectivore (small insects)	4.64	0.68	3.43	0.50	2.74	0.40	0.28	0.04	2.59	0.38	1.91	0.28	1.53	0.22	0.16	0.02
	6.8	Insectivore (large insects)	1.16	0.17	0.86	0.13	0.68	0.10	0.07	0.01	0.55	0.08	0.41	0.06	0.33	0.05	0.03	0.00
	6.8	Granivore (grain and seeds)	1.16	0.17	0.86	0.13	0.68	0.10	0.07	0.01	0.55	0.08	0.41	0.06	0.33	0.05	0.03	0.00
Reproduction	6.8	Frugivore (fruit)	2.32	0.34	1.72	0.25	1.37	0.20	0.14	0.02	1.11	0.16	0.82	0.12	0.65	0.10	0.07	0.01
	6.8	Herbivore (short grass)	16.57	2.44	12.26	1.80	9.78	1.44	0.99	0.15	5.89	0.87	4.36	0.64	3.47	0.51	0.35	0.05
	6.8	Herbivore (long grass)	10.12	1.49	7.49	1.10	5.97	0.88	0.61	0.09	3.30	0.49	2.45	0.36	1.95	0.29	0.20	0.03
	6.8	Herbivore (forage crops)	15.33	2.25	11.35	1.67	9.05	1.33	0.92	0.14	5.07	0.75	3.75	0.55	2.99	0.44	0.30	0.04

Table 33 Screening Level Risk Assessment to Mammals (Luna Privilege)

Effects	Toxicity (mg a.i./kg bw/d)	Feeding Guild (food item)	EDE (mg a.i./kg bw)	RQ
	5	Small Mammal (0.015 kg)		
Acute	200.00*	Insectivore (small insects)	11.71	0.06
Reproduction	13.90	Insectivore (small insects)	11.71	0.84
	Medi	um Sized Mammal (0.035 kg)		
Acute	200.00*	Herbivore (short grass)	36.67	0.18
Reproduction	13.90	Herbivore (short grass)	36.67	2.64
	L	arge Sized Mammal (1 kg)		
Acute	200.00*	Herbivore (short grass)	19.60	0.10
Reproduction	13.90	Herbivore (short grass)	19.60	1.41

*based on an uncertainty factor of 10

Table 34 Expanded Screening Level Assessment of Reproductive Risk to Mammals with Same Endpoints (Luna Privilege)

Effects	Toxicity	Food Guild (food item)			Max	imun	Resid	lues					M	ean F	tesidu	es		
	(mg a.i./kg bw/d)		On-	field	Off 1		Off I		Off (6	Field %)	On-	field	Off 1	Field %)	Off 1		Off (6°	Field %)
			EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ
			1	Smal	l Man	amal	(0.015	kg)		_		_				_		_
Reproduction	13.90	Insectivore (small insects)	11.71	0.84	8.66	0.62	6.91	0.50	0.70	0.05	6.53	0.47	4.83	0.35	3.85	0.28	0.39	0.03
	13.90	Granivore (grain and seeds)	2.93	0.21	2.17	0.16	1.73	0.12	0.18	0.01	1.40	0.10	1.03	0.07	0.82	0.06	0.08	0.01
	13.90	Frugivore (fruit)	5.85	0.42	4.33	0.31	3.45	0.25	0.35	0.03	2.79	0.20	2.07	0.15	1.65	0.12	0.17	0.01
			Med	lium	Sized	Mam	mal (0	.035	kg)	_				_		_		_
Reproduction	13.90	Insectivore (small insects)	10.26	0.74	7.59	0.55	6.05	0.44	0.62	0.04	5.72	0.41	4.23	0.30	3.38	0.24	0.34	0.02
	13.90	Insectivore (large insects)	2.57	0.18	1.90	0.14	1.51	0.11	0.15	0.01	1.22	0.09	0.91	0.07	0.72	0.05	0.07	0.01
	13.90	Granivore (grain and seeds)	2.57	0.18	1.90	0.14	1.51	0.11	0.15	0.01	1.22	0.09	0.91	0.07	0.72	0.05	0.07	0.01
	13.90	Frugivore (fruit)	5.13	0.37	3.80	0.27	3.03	0.22	0.31	0.02	2.45	0.18	1.81	0.13	1.44	0.10	0.15	0.01
	13.90	Herbivore (short grass)	36.67	2.64	27.14	1.95	21.64	1.56	2.20	0.16	13.02	0.94	9.64	0.69	7.68	0.55	0.78	0.06
	13.90	Herbivore (long grass)	22.39	1.61	16.57	1.19	13.21	0.95	1.34	0.10	7.31	0.53	5.41	0.39	4.31	0.31	0.44	0.03
	13.90	Herbivore (forage crops)	33.93	2.44	25.11	1.81	20.02	1.44	2.04	0.15	11.22	0.81	8.30	0.60	6.62	0.48	0.67	0.05
			1	arge	Sized	Man	mal (l kg)										
Reproduction	13.90	Insectivore (small insects)	5.48	0.39	4.06	0.29	3.24	0.23	0.33	0.02	3.06	0.22	2.26	0.16	1.80	0.13	0.18	0.01
	13.90	Insectivore (large insects)	1.37	0.10	1.01	0.07	0.81	0.06	0.08	0.01	0.65	0.05	0.48	0.03	0.39	0.03	0.04	0.00
	13.90	Granivore (grain and seeds)	1.37	0.10	1.01	0.07	0.81	0.06	0.08	0.01	0.65	0.05	0.48	0.03	0.39	0.03	0.04	0.00
	13.90	Frugivore (fruit)	2.74	0.20	2.03	0.15	1.62	0.12	0.16	0.01	1.31	0.09	0.97	0.07	0.77	0.06	0.08	0.01
	13.90	Herbivore (short grass)	19.60	1.41	14.50	1.04	11.56	0.83	1.18	0.08	6.96	0.50	5.15	0.37	4.11	0.30	0.42	0.03
	13.90	Herbivore (long grass)	11.97	0.86	8.85	0.64	7.06	0.51	0.72	0.05	3.91	0.28	2.89	0.21	2.31	0.17	0.23	0.02
	13.90	Herbivore (forage crops)	18.13	1.30	13.42	0.97	10.70	0.77	1.09	0.08	5.99	0.43	4.44	0.32	3.54	0.25	0.36	0.03

Table 35 Refined Assessment of Reproductive Risk to Mammals (Luna Privilege)

	Toxicity				Max	cimu	m Resi	dues					N	lean	Resid	ues		
Effects	(mg a.i./kg	Food Guild (food item)	On-	field	Off I		Off 1 (59			Field %)	On-1	ield	Off 1			Field 9%)		Field %)
	bw/d)		EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ
Small Mamm	al (0.015 k	g)				-												
	82.4	Insectivore (small insects)	11.71	0.14	8.66	0.11	6.91	0.08	0.70	0.01	6.53	0.08	4.83	0.06	3.85	0.05	0.39	< 0.01
Reproduction	82.4	Granivore (grain and seeds)	2.93	0.04	2.17	0.03	1.73	0.02	0.18	< 0.01	1.40	0.02	1.03	0.01	0.82	0.01	80.0	< 0.01
	82.4	Frugivore (fruit)	5.85	0.07	4.33	0.05	3.45	0.04	0.35	< 0.01	2.79	0.03	2.07	0.03	1.65	0.02	0.17	< 0.01
Medium Size	d Mamma	(0.035 kg)																
	82.4	Insectivore (small insects)	10.26	0.12	7.59	0.09	6.05	0.07	0.62	0.01	5.72	0.07	4.23	0.05	3.38	0.04	0.34	<0.01
	82.4	Insectivore (large insects)	2.57	0.03	1.90	0.02	1.51	0.02	0.15	< 0.01	1.22	0.01	0.91	0.01	0.72	0.01	0.07	< 0.01
	82.4	Granivore (grain and seeds)	2.57	0.03	1.90	0.02	1.51	0.02	0.15	< 0.01	1.22	0.01	0.91	0.01	0.72	0.01	0.07	< 0.01
Reproduction	82.4	Frugivore (fruit)	5.13	0.06	3.80	0.05	3.03	0.04	0.31	< 0.01	2.45	0.03	1.81	0.02	1.44	0.02	0.15	< 0.01
	82.4	Herbivore (short grass)	36.67	0.45	27.14	0.33	21.64	0.26	2.20	0.03	13.02	0.16	9.64	0.12	7.68	0.09	0.78	0.01
	82.4	Herbivore (long grass)	22.39	0.27	16.57	0.20	13.21	0.16	1.34	0.02	7.31	0.09	5.41	0.07	4.31	0.05	0.44	0.01
	82.4	Herbivore (forage crops)	33.93	0.41	25.11	0.30	20.02	0.24	2.04	0.02	11.22	0.14	8.30	0.10	6.62	0.08	0.67	0.01
Large Sized N	Jammal (1	kg)																
	82.4	Insectivore (small insects)	5.48	0.07	4.06	0.05	3.24	0.04	0.33	< 0.01	3.06	0.04	2.26	0.03	1.80	0.02	0.18	< 0.01
Reproduction	82.4	Insectivore (large insects)	1.37	0.02	1.01	0.01	0.81	0.01	0.08	< 0.01	0.65	0.01	0.48	0.01	0.39	< 0.01	0.04	< 0.01

	Toxicity				Max	imui	n Resi	dues					N	Jean	Resid	ues		
Effects	(mg a.i./kg	Food Guild (food item)	On-field		-field Off Fig. (74%)		Off I		1	Field %)	On-f	ield	Off 1	Field %)	-	Field (%)		Field %)
	bw/d)		EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ
	82.4	Granivore (grain and seeds)	1.37	0.02	1.01	0.01	0.81	0.01	0.08	< 0.01	0.65	0.01	0.48	0.01	0.39	<0.01	0.04	< 0.0
	82.4	Frugivore (fruit)	2.74	0.03	2.03	0.02	1.62	0.02	0.16	< 0.01	1.31	0.02	0.97	0.01	0.77	0.01	0.08	< 0.01
	82.4	Herbivore (short grass)	19.60	0.24	14.50	0.18	11.56	0.14	1.18	0.01	6.96	0.08	5.15	0.06	4.11	0.05	0.42	0.01
	82.4	Herbivore (long grass)	11.97	0.15	8.85	0.11	7.06	0.09	0.72	0.01	3.91	0.05	2.89	0.04	2.31	0.03	0.23	< 0.01
	82.4	Herbivore (forage crops)	18.13	0.22	13.42	0.16	10.70	0.13	1.09	0.01	5.99	0.07	4.44	0.05	3.54	0.04	0.36	< 0.01

Table 36 Screening Level Risk Assessment to Terrestrial Plants (Luna Privilege)

Organism	Exposure	Test Substance	Tox Value for RQ	EEC	RQ
Vascular plants	seedling emergence	Luna Privilege A G	EC ₂₅ : 500 g a.i./ha	497.76 g a.i./ ha	1.00
	vegetative vigour	Luna Privilege A G	EC ₂₅ : 250 g a.i./ha	403.9 g a.i./ha*	1.62

*with a default foliar half-life of 10 days

Table 37 Refined Risk Assessment to Terrestrial Plants (Luna Privilege)

	Airblast Early (74% drift)	Airblast Late (59% drift)	Ground Boom (6% drift)
Application rate (250 g a.i./ha)	185 g a.i./ha	147.5 g a.i./ha	15.00 g a.i/ha
Seedling Er	mergence		
Cumulative application rate (2 applications, 7 d interval and DT50 of 539 days)	368.34 g a.i./ha	293.68 g a.i./ha	29.87 g a.i./ha
RQ with EC ₂₅ of 500 g a.i./ha for seedling emergence	0.77	0.59	0.06
Risk	no risk	no risk	no risk
Vegitative	vigour		
Cumulative application rate (2 applications, 7 d interval and foliar half-life of 10 days)	298.89 g a.i./ha	238.31 g a.i./ha	24.24 g a.i./ha
RQ with EC ₂₅ of 250 g a.i./ha for vegitative vogour	1.2	0.95	0.1
Risk	risk	no risk	no risk

Table 38 Screening Level EEC* for Fluopyram (Luna Tranquility Fungicide)

Soil**	Wate	r***
	15 cm depth	80 cm depth
0.22 mg a.i./kg soil	0.33 mg a.i./L water	0.33 mg a.i./L water

*based on 5 applications of 100 g a.i./ha each with a cumulative application rate 491.12 g

**top 30 cm soil depth and a soil bulk density of 1.5 g/cm³

*** cumulative application of 496.72 g a.i./ha based on a half-life of 1470 days

Table 39 Maximum EECs Fluopyram in Vegetation and Insects after a Direct Over-Spray (Luna Tranquility Fungicide)

Matrix	EEC ^a (mg a.i./kg fw)	Fresh/Dry Weight Ratios	EEC Direct Overspray (mg a.i./kg dw) Maximum Residues	Mean EEC (mg a.i./kg dw) Mean Residues
Short range grass	50.7537	3.3	167.4873	59.4816
Leaves and leafy crops	28.6966	11	315.6628	104.3513
Long grass	23.2418	4,4	102.2641	33.3924
Forage crops	28.6966	5.4	154.9617	51.2270
Small insects	12.3324	3.8	46.8631	26.1352
Pods with seeds	3.0831	3.9	12.0242	5.7346
Large insects	3.0831	3.8	11.7158	5.5876
Grain and seeds	3.0831	3.8	11.7158	5.5876
Fruit	3.0831	7.6	23.4317	11.1751

^a Cumulative application rate of 237.162 g a.i./ha (five applications of 100 g a.i. each with 7-day interval and with default half-life of 10 days)

Table 40 Screening Level Risk Assessment to Wild Birds (Luna Tranquility Fungicide)

Effects	Toxicity (mg a.i./kg bw/d)	Feeding Guild (food item)	EDE (mg a.i./kg bw)	RQ
120000000000000000000000000000000000000	S	Small Bird (0.02 kg)		
Acute	200.00	Insectivore (small insects)	11.95	0.06
Reproduction	4.12	Insectivore (small insects)	11.95	2.90
	Med	ium Sized Bird (0.1 kg)		
Acute	200.00	Insectivore (small insects)	9.33	0.05
Reproduction	4.12	Insectivore (small insects)	9.33	2.26
	La	arge Sized Bird (1 kg)		
Acute	200.00	Herbivore (short grass)	9.73	0.05
Reproduction	4.12	Herbivore (short grass)	9.73	2.36

Table 41 Expanded Screening Level Reproductive Risk Assessment for Wild Birds for On-Field and Off-Field Scenarios (Luna Tranquility Fungicide)

	Toxicity				Max	imur	n Resid	lues					M	ean B	tesidu	es		
Effects	(mg a.i./kg	Food Guild (food item)	On-	field	Off-field (74%)		Off-1		Off-		On-1	field	Off 1		Off 1 (59			Field %)
	bw/d)		EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ
				S	mall l	Bird (0.02 kg	()								and the second		
	4.12	Insectivore (small insects)	11.95	2.90	8.84	2.15	7.05	1.71	0.72	0.17	6.66	1.62	4.93	1.20	3.93	0.95	0.40	0.10
Reproduction	4.12	Granivore (grain and seeds)	2.99	0.73	2.21	0.54	1.76	0.43	0.18	0.04	1.42	0.35	1.05	0.26	0.84	0.20	0.09	0.02
	4.12	Frugivore (fruit)	5.98	1.45	4.42	1.07	3.53	0.86	0.36	0.09	2.85	0.69	2.11	0.51	1.68	0.41	0.17	0.04
				Medi	um Si	zed E	ird (0.	1 kg)										
D di		Insectivore (small insects)	9.33	2.26	6.90	1.68	5.50	1.34	0.56	0.14	5.20	1.26	3.85	0.93	3.07	0.74	0.31	0.08
Reproduction	4.12	Insectivore (large insects)	2.33	0.57	1.73	0.42	1.38	0.33	0.14	0.03	1.11	0.27	0.82	0.20	0.66	0.16	0.07	0.02

	Toxicity				Max	imur	n Resid	lues					M	ean F	tesidu	es		
Effects	(mg a.i./kg	Food Guild (food item)	On-	field	Off-	-	Off-6		Off-1		On-f	ield	Off 1		Off I		Off 1	
	bw/d)		EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ
	4.12	Granivore (grain and seeds)	2.33	0.57	1.73	0.42	1.38	0.33	0.14	0.03	1.11	0.27	0.82	0.20	0.66	0.16	0.07	0.02
	4.12	Frugivore (fruit)	4.66	1.13	3.45	0.84	2.75	0.67	0.28	0.07	2.22	0.54	1.65	0.40	1.31	0.32	0.13	0.03
				La	rge Si	zed B	ird (1 l	kg)										
	4.12	Insectivore (small insects)	2.72	0.66	2.01	0.49	1.61	0.39	0.16	0.04	1.52	0.37	1.12	0.27	0.90	0.22	0.09	0.02
	4.12	Insectivore (large insects)	0.68	0.17	0.50	0.12	0.40	0.10	0.04	0.01	0.32	0.08	0.24	0.06	0.19	0.05	0.02	0.00
	4.12	Granivore (grain and seeds)	0.68	0.17	0.50	0.12	0.40	0.10	0.04	0.01	0.32	0.08	0.24	0.06	0.19	0.05	0.02	0.00
Reproduction	4.12	Frugivore (fruit)	1.36	0.33	1.01	0.24	0.80	0.19	0.08	0.02	0.65	0.16	0.48	0.12	0.38	0.09	0.04	0.01
	4.12	Herbivore (short grass)	9.73	2.36	7.20	1.75	5.74	1.39	0.58	0.14	3.46	0.84	2.56	0.62	2.04	0.49	0.21	0.05
	4.12	Herbivore (long grass)	5.94	1.44	4.40	1.07	3.51	0.85	0.36	0.09	1.94	0.47	1.44	0.35	1.14	0.28	0.12	0.03
	4.12	Herbivore (forage crops)	9.00	2.19	6.66	1.62	5.31	1.29	0.54	0.13	2.98	0.72	2.20	0.53	1.76	0.43	0.18	0.04

Table 42 Refined Assessment of Reproductive Risk for Wild Birds (Luna Tranquility Fungicide)

	Toxicity				Max	imun	Resid	dues					M	ean R	tesidu	es		
Effects	(mg a.i./kg	Food Guild (food item)	On-	field		field %)	Off-		Off-		On-	field	Off 1	Field %)	Off 1 (59		Off I	
	bw/d)		EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ
				Si	mall E	Bird ().02 k	g)					-					
	6.8	Insectivore (small insects)	11.95	1.76	8.84	1.30	7.05	1.04	0.72	0.11	6.66	0.98	4.93	0.73	3.93	0.58	0.40	0.06
Reproduction	6.8	Granivore (grain and seeds)	2.99	0.44	2.21	0.33	1.76	0.26	0.18	0.03	1.42	0.21	1.05	0.16	0.84	0.12	0.09	0.01
	6.8	Frugivore (fruit)	5.98	0.88	4.42	0.65	3.53	0.52	0.36	0.05	2.85	0.42	2.11	0.31	1.68	0.25	0.17	0.03
				Medi	um Si	zed B	ird (0.	.1 kg)										
	6.8	Insectivore (small insects)	9.33	1.37	6.90	1.01	5.50	0.81	0.56	0,08	5.20	0.76	3.85	0.57	3.07	0.45	0.31	0.05
n 1 4	6.8	Insectivore (large insects)	2.33	0.34	1.73	0.25	1.38	0.20	0.14	0.02	1.11	0.16	0.82	0.12	0.66	0.10	0.07	0.01
Reproduction	6.8	Granivore (grain and seeds)	2.33	0.34	1.73	0.25	1.38	0.20	0.14	0.02	1.11	0.16	0.82	0.12	0.66	0.10	0.07	0.01
	6.8	Frugivore (fruit)	4.66	0.69	3.45	0.51	2.75	0.40	0.28	0.04	2.22	0.33	1.65	0.24	1.31	0.19	0.13	0.02
				Lai	rge Siz	zed Bi	ird (1	kg)										
	6.8	Insectivore (small insects)	2.72	0.40	2.01	0.30	1.61	0.24	0.16	0.02	1.52	0.22	1.12	0.17	0.90	0.13	0.09	0.01
	6.8	Insectivore (large insects)	0.68	0.10	0.50	0.07	0.40	0.06	0.04	0.01	0.32	0.05	0.24	0.04	0.19	0.03	0.02	0.00
	6.8	Granivore (grain and seeds)	0.68	0.10	0.50	0.07	0.40	0.06	0.04	0.01	0.32	0.05	0.24	0.04	0.19	0.03	0.02	0.00
Reproduction	6.8	Frugivore (fruit)	1.36	0.20	1.01	0.15	0.80	0.12	0.08	0.01	0.65	0.10	0.48	0.07	0.38	0.06	0.04	0.01
	6.8	Herbivore (short grass)	9.73	1.43	7.20	1.06	5.74	0.84	0.58	0.09	3.46	0.51	2.56	0.38	2.04	0.30	0.21	0.03
	6.8	Herbivore (long grass)	5.94	0.87	4.40	0.65	3.51	0.52	0.36	0.05	1.94	0.29	1.44	0.21	1.14	0.17	0.12	0.02
	6.8	Herbivore (forage crops)	9.00	1.32	6.66	0.98	5.31	0.78	0.54	0.08	2.98	0.44	2.20	0.32	1.76	0.26	0.18	0.03

Table 43 Screening Level Risk Assessment to Mammals (Luna Tranquility Fungicide)

Effects	Toxicity (mg a.i./kg bw/d)	Feeding Guild (food item)	EDE (mg a.i./kg bw)	RQ
	Small	Mammal (0.015 kg)		
Acute	200.00	Insectivore (small insects)	6.87	0.03
Reproduction	13.90	Insectivore (small insects)	6.87	0.49
1	Medium Sized Mammal	(0.035 kg)		
Acute	200.00	Herbivore (short grass)	21.53	0.11
Reproduction	13.90	Herbivore (short grass)	21.53	1.55
	Large	Sized Mammal (1 kg)		
Acute	200.00	Herbivore (short grass)	11.51	0.06
Reproduction	13.90	Herbivore (short grass)	11.51	0.83

Table 44 Expanded Screening Level Reproductive Risk Assessment for Mammals (Luna Tranquility Fungicide)

	Toxicity				Maxi	mum	Resid	lues					M	ean F	Residu	es		
Effects	(mg a.i./kg	Food Guild (food item)	On-f	ield	Off I		Off I		Off 1		On-f	field	Off 1		Off I		Off 1	
	bw/d)		EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ
			N	lediu	ım Ma	mma	1 (0.05	5 kg		-					-			
	13.90	Insectivore (small insects)	6.03	0.43	4.46	0.32	3.55	0.26	0.36	0.03	3.36	0.24	2.49	0.18	1.98	0.14	0.20	0.01
	13.90	Insectivore (large insects)	1.51	0.11	1.11	0.08	0.89	0.06	0.09	0.01	0.72	0.05	0.53	0.04	0.42	0.03	0.04	0.00
	13.90	Granivore (grain and seeds)	1.51	0.11	1.11	0.08	0.89	0.06	0.09	0.01	0.72	0.05	0.53	0.04	0.42	0.03	0.04	0.00
Reproduction	13.90	Frugivore (fruit)	3.01	0.22	2.23	0.16	1.78	0.13	0.18	0.01	1.44	0.10	1.06	0.08	0.85	0.06	0.09	0.01
	13.90	Herbivore (short grass)	21.53	1.55	15.94	1.15	12.71	0.91	1.29	0.09	7.65	0.55	5.66	0.41	4.51	0.32	0.46	0.03
	13.90	Herbivore (long grass)	13.15	0.95	9.73	0.70	7.76	0.56	0.79	0.06	4.29	0.31	3.18	0.23	2.53	0.18	0.26	0.02
	13.90	Herbivore (forage crops)	19.92	1.43	14.74	1.06	11.75	0.85	1.20	0.09	6.59	0.47	4.87	0.35	3.89	0.28	0.40	0.03

Table 45 Refined Assessment of Reproductive Risk for Mammals (Luna Tranquility Fungicide)

	Toxicity				Max	imun	Resid	lues					M	ean I	Residu	es		
Effects	(mg a.i./kg	Food Guild (food item)	On-	lield	Off 1		Off I		Off 1		On-	field	Off 1		Off 1		Off 1	
	bw/d)		EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ
			Me	dium	sized	Mam	mal (0	.035	kg)					-				
	82.4	Insectivore (small insects)	6.03	0.07	4.46	0.05	3.55	0.04	0.36	0.00	3.36	0.04	2.49	0.03	0.04	1.98	0.20	0.00
	82.4	Insectivore (large insects)	1.51	0.02	1.11	0.01	0.89	0.01	0.09	0.00	0.72	0.01	0.53	0.01	0.01	0.42	0.04	0.00
	82.4	Granivore (grain and seeds)	1.51	0.02	1.11	0.01	0.89	0.01	0.09	0.00	0.72	0.01	0.53	0.01	0.01	0.42	0.04	0.00
Reproduction	82.4	Frugivore (fruit)	3.01	0.04	2.23	0.03	1.78	0.02	0.18	0.00	1.44	0.02	1.06	0.01	0.02	0.85	0.09	0.00
	82.4	Herbivore (short grass)	21.53	0.26	15.94	0.19	12.71	0.15	1.29	0.02	7.65	0.09	5.66	0.07	0.09	4.51	0.46	0.01
	82.4	Herbivore (long grass)	13.15	0.16	9.73	0.12	7.76	0.09	0.79	0.01	4.29	0.05	3.18	0.04	0.05	2.53	0.26	0.00
	82.4	Herbivore (forage crops)	19.92	0.24	14.74	0.18	11.75	0.14	1.20	0.01	6.59	0.08	4.87	0.06	0.08	3.89	0.40	0.00

Table 46 Screening Level Risk Assessment to Terrestrial Plants (Luna Tranquility Fungicide)

Organism	Exposure	Test Substance	Tox Value For RQ	EEC	RQ
Vascular plants	seedling emergence	AE C656948 SC 500A G	EC ₂₅ : 500 g a.i./ha	491.12 g a.i./ ha*	0.98
	vegetative vigour	AE C656948 SC 500A G	EC ₂₅ : 250 g a.i./ha	237.162 g a.i./ha	0.95

^{*}based the cumulative rate with a field DT50 of 539 days; ** with a default foliar half-life of 10 days

Table 47 Screening Level Risk Assessment to Wild Birds (Propulse Fungicide)

Effects	Toxicity (mg a.i./kg bw/d)	Feeding Guild (food item)	EDE (mg a.i./kg bw)	RQ
		Small Bird (0.02 kg)		
Acute	200.00	Insectivore (small insects)	12.21	0.06
Reproduction	4.12	Insectivore (small insects)	12.21	2.96
	Me	edium Sized Bird (0.1 kg)		
Acute	200.00	Insectivore (small insects)	9.53	0.05
Reproduction	4.12	Insectivore (small insects)	9.53	2.31
		Large Sized Bird (1 kg)		
Acute	200.00	Herbivore (short grass)	9.94	0.05
Reproduction	4.12	Herbivore (short grass)	9.94	2.41

Table 48 Expanded Screening Level Reproductive Risk Assessment for Wild Birds (Propulse Fungicide)

	Toxicity		N	1axim	um resi	dues		Mea	n residu	es
Effects	mg	Food Guild (food item)	On-	field	Off-fi	eld (6%)	On-	-field	Off Fie	dd (6%)
Effects	a.i./kg bw/d	Toou Guila (roou item)	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ
		Small	Bird (0.	02 kg)						
	4.12	Insectivore (small insects)	12.21	2.96	0.73	0.18	6.81	1.65	0.41	0.10
Reproduction	4.12	Granivore (grain and seeds)	3.05	0.74	0.18	0.04	1.46	0.35	0.09	0.02
	4.12	Frugivore (fruit)	6.11	1.48	0.37	0.09	2.91	0.71	0.17	0.04
		Medium S	ized Bi	rd (0.1	kg)			•		
	4.12	Insectivore (small insects)	9.53	2.31	0.57	0.14	5.31	1.29	0.32	0.08
D 1 1	4.12	Insectivore (large insects)	2.38	0.58	0.14	0.03	1.14	0.28	0.07	0.02
Reproduction	4.12	Granivore (grain and seeds)	2.38	0.58	0.14	0.03	1.14	0.28	0.07	0.02
	4.12	Frugivore (fruit)	4.76	1.16	0.29	0.07	2.27	0.55	0.14	0.03

	Toxicity		N	Taximu	um resid	dices		Mea	n residu	es
Effects	mg	Food Guild (food item)	On-	field	Off-in	eld (6%)	On-	field	Off Fie	dd (6%)
Effects	a.i./kg bw/d	rood Gana (tood item)	EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ
		Large S	ized Bir	d (1 kg	g)					
	4.12	Insectivore (small insects)	2.78	0.68	0.17	0.04	1.55	0.38	0.09	0.02
	4.12	Insectivore (large insects)	0.70	0.17	0.04	0.01	0.33	0.08	0.02	0.00
	4.12	Granivore (grain and seeds)	0.70	0.17	0.04	0.01	0.33	0.08	0.02	0.00
Reproduction	4.12	Frugivore (fruit)	1.39	0.34	0.08	0.02	0.66	0.16	0.04	0.01
	4.12	Herbivore (short grass)	9.94	2.41	0.60	0.14	3.53	0.86	0.21	0.05
	4.12	Herbivore (long grass)	6.07	1.47	0.36	0.09	1.98	0.48	0.12	0.03
	4.12	Herbivore (forage crops)	9.20	2.23	0.55	0.13	3.04	0.74	0.18	0.04

Table 49 Refined Reproductive Risk Assessment for Wild Birds (Propulse Fungicide)

			N	laxim	um resi	lues		Mea	n residue	28
Effects	Toxicity	Food Guild (food item)	On-	field	Off-fi	eld (6%)	On-	field	Off Fie	ld (6%)
			EDE	RQ	EDE	RQ	EDE	RQ	EDE	RQ
		Small	Bird (0.	.02 kg))					
	6.8	Insectivore (small insects)	12.21	1.80	0.73	0.11	6.81	1.00	0.41	0.06
Reproduction	6.8	Granivore (grain and seeds)	3.05	0.45	0.18	0.03	1.46	0.21	0.09	0.01
	6.8	Frugivore (fruit)	6.11	0.90	0.37	0.05	2.91	0.43	0.17	0.03
		Medium S	ized Bi	rd (0.1	kg)					
	6.8	Insectivore (small insects)	9.53	1.40	0.57	0.08	5.31	0.78	0.32	0.05
D 1 1	6.8	Insectivore (large insects)	2.38	0.35	0.14	0.02	1.14	0.17	0.07	0.01
Reproduction	6.8	Granivore (grain and seeds)	2.38	0.35	0.14	0.02	1.14	0.17	0.07	0.01
	6.8	Frugivore (fruit)	4.76	0.70	0.29	0.04	2.27	0.33	0.14	0.02
		Large Si	ized Bir	rd (1 k	g)					
	6.8	Insectivore (small insects)	2.78	0.41	0.17	0.02	1.55	0.23	0.09	0.01
	6.8	Insectivore (large insects)	0.70	0.10	0.04	0.01	0.33	0.05	0.02	0.00
	6.8	Granivore (grain and seeds)	0.70	0.10	0.04	0.01	0.33	0.05	0.02	0.00
Reproduction	6.8	Frugivore (fruit)	1.39	0.20	0.08	0.01	0.66	0.10	0.04	0.01
	6.8	Herbivore (short grass)	9.94	1.46	0.60	0.09	3.53	0.52	0.21	0.03
	6.8	Herbivore (long grass)	6.07	0.89	0.36	0.05	1.98	0.29	0.12	0.02
	6.8	Herbivore (forage crops)	9.20	1.35	0.55	0.08	3.04	0.45	0.18	0.03

Table 50 Screening Level Risk Assessment to Terrestrial Plants (Propulse Fungicide)

Organism	Exposure	Test Substance	Tox Value For RQ	EEC	RQ
Vascular plants	seedling emergence	AE C656948 SC 500A G	EC25: 500.00 g a.i./ha	298.656 g a.i./ ha	0.60
,	vegetative vigour	AE C656948 SC 500A G	EC25:250.00g a.i./ha	242.345 g a.i./ha	0.97

Table 51 Effects on Aquatic Organisms

Organism	Exposure	Test Substance	Endpoint Value	Degree of Toxicity	PMRA#
		Freshwater	species		
Rainbow trout (Oncorhynchus	acute (96 h)		LC ₅₀ : >1.78 mg a.i./L NOAEC: 1.78 mg a.i./L	moderately toxic	1599539
mykiss)	acute (96 h)	Luna Privilege G	LC ₅₀ : >46.4 mg a.i./L NOAEC: 1.31 mg a.i./L EC ₅₀ : 3.71 mg a.i./L (sub-lethal effects)	slightly toxic	
Bluegill sunfish (Lepomis macrochirus)	acute (96 h)	AE C656948	LC ₅₀ : >5.17 mg a.i./L NOAEC: 5.17 mg a.i./L	moderately toxic	1599538
Fathead minnow (Pimephales promelas)	acute (96 h)	AE C656948	LC ₅₀ : >4.95 mg a.i./L NOAEC: 4.95 mg a.i./L EC ₅₀ : >4.95 mg a.i./L	moderately toxic	1599543
Fathcad minnow (Pimephales promelas)	chronic: early life stage (33 d)	AE C656948	NOAEC: 0.135 mg a.i./L LOAEC: 0.269 mg a.i./L	N/A	1599730
Daphnia (Daphnia magna)	acute (48 h)	AE C656948	NOAEC: 17 mg a.i./L EC ₅₀ : >17 mg a.i./L	slightly toxic	1599541
	chronic (Life cycle: 21 d)	AE C656948	NOAEC: 1.214 mg a.i./L LOAEC: 2.996 mg a.i./L 21-d EC ₅₀ : 2.700 mg a.i./L (reproduction)	N/A	1599770
	acute (48 h)	Luna Privilege G	48-h EC ₅₀ ; >38.2 mg a.i./L NOAEC: 11.6 mg a.i./L	slightly toxic	1599285
Freshwater green algae	acute (96 h)	AE C656948	NOAEC: 1.46 mg a.i./L EC ₅₀ : 4.3 mg a.i./L (Biomass)	N/A	1599864
(Pseudokirchneriella subcapitata)	acute (72 h)	Luna Privilege G	NOAEC: 1.17 mg a.i./L EC ₀₅ : 1.0 mg a.i./L EC ₅₀ : 3.4 mg a.i./L (Cell density)	N/A	1599287
	acute (72 h)	Fluopyram- Lactame (a metabolite of Fluopyram)	NOAEC: 8.87 mg a.i./L EC ₅₀ : >8.87 mg a.i./L (growth	N/A	1599808
Freshwater diatom (Navicula pelliculosa)	acute (96 h)		NOAEC: 2.47 mg a.i./L EC ₅₀ : 6.1 mg a.i./L (biomass)	N/A	1599862
Freshwater blue- green algae (Anabaena flos- aquae)	acute (96 h)	AE C656948	NOAEC: 9.69 mg a.i./L EC ₅₀ : >9.69 mg a.i./L most sensitive end-point: none	N/A	1599863
	acute (7 d)	AE C656948	NOAEC: 0.278 mg a.i./L EC ₀₅ > 0.278 mg a.i./L EC ₅₀ : 2.6 mg a.i./L (frond number based on yield)	N/A	1599773
	acute (7 d)	Luna Privilege G	NOAEC: 1.04 mg a.i./L EC ₀₅ : 1.8 mg a.i./L EC ₅₀ : 2.9 mg a.i./L (frond number)	N/A	1599303
Sediment dwelling Freshwater chironomid (Chironomus riparius)	28-d chronic	AE C656948	overlying water concentrations* EC ₅₀ (emergence ratio): >5.52 mg a.i./L NOAEC (emergence ratio): >0.0128 and <3.11 mg a.i./L	N/A	1599633

Organism	Exposure	Test Substance	Endpoint Value	Degree of Toxicity	PMRA#
			0.525 mg a.i./L (TWA)		
Freshwater dipteran midge (Chironomus tentans)	54-day life-cycle	AE C656948	NOAEC (survival and emergences): sediment: 26 mg a.i./kg pore water: 3.8 mg a.i./L (TWA) overlying water: 0.14 mg a.i./L	N/A	1599614
Marine species	•	1			
Sheepshead Minnow (Cyprinodon variegatus)	acute (96 h)	AE C656948	LC ₅₀ : >0.98 mg a.i./L NOAEC: 0.98 mg a.i./L EC ₅₀ : >0.98 mg a.i./L	highly toxic	1599544
Eastern Oyster (Crassostrea virginica)	acute (96 h)	AE C656948	EC ₅₀ : >0.43 mg a.i./L NOAEC: 0.43 mg a.i./L (shell deposition)	highly toxic	1599604
Saltwater Mysid (Americamysis bahia)	acute (96 h)	AE C656948	LC ₅₀ : >0.51 mg a.i./L NOAEC: 0.27 mg a.i./L	highly toxic	1599603
Saltwater Diatom (Skeletonema costatum)	acute (96 h)	AE C656948 EC ₅₀ : > 1.13 mg a.i./L mode NOAEC: 1.13 mg a.i./L (cell density, biomass, growth rate) most sensitive end-point: none			1599865
Marine amphipods (Leptocheirus plumulosus)	10-d acute	AE C656948 sediment concentrations LC ₅₀ mortality: >100 mg a.i./kg NOAEC (mortality): 100 mg a.i./kg Pore water concentrations LC ₅₀ mortality: >7.5 mg a.i./L NOAEC (mortality): 7.5 mg a.i./L Overlying water concentrations LC ₅₀ mortality: >1.6 mg a.i./L NOAEC (mortality): 1.6 mg a.i./L		moderately toxic	1599616
Marine amphipods (Leptocheirus plumulosus)	28-d chronic	AE C656948	Rediment concentrations (Total radioactive residues equivalent to a.i) EC ₅₀ growth: >92 mg a.i./kg NOAEC (growth): 36 mg a.i./kg pore water concentrations EC ₅₀ growth: >5.9 mg a.i./L NOAEC (growth): 2.5 mg a.i./L overlying water concentrations EC ₅₀ growth: >1.19 mg a.i./L NOAEC (growth): 0.55 mg a.i./L		1599615

^{*}sediment concentrations not measured; **TWA Time weighted average

Table 52 Screening Level Risk Assessment to Aquatic Organisms (Luna Privilege)

Organism	Exposure	Test Substance	Tox Value For RQ (mg a.i./L)	EEC (mg a.i./L)	RQ
Rainbow trout (O. mykiss)	acute	AEC 656948	(LC ₅₀ /10): >0.178**	0.062*	< 0.35
Bluegill sunfish (L. macrochirus)	bioaccumulation study	AE C656948	BCF: 18	low potential for bioaccumulation	
Fathead minnow (P. promelas)	chronic: early life stage	AE C656948	NOAEC: 0.135	0.062*	0.46
Sediment dwelling	54 d pore water	AE C656948	NOAEC: 3.8	0.062*	0.16
(C. riparius) (C. tetans)	54 d sediment	AE C656948	NOAEC: 26	0.062*	0.002
Amphibians	acute	AE C656948	(LC ₅₀ /10): >0.178**	0.33 [£]	<1.85
	chronic	AE C656948	NOAEC: 0.135	0.33 [£]	2.44
Daphnia	Acute	AE C656948	(EC ₅₀ /2): >8.5***	0.062*	< 0.017
(D. magna)	chronic	AE C656948	NOAEC: 1.214	0.062*	0.05
Freshwater green algae (<i>P. subcapitata</i>)	acute	Luna Privilege G	(EC ₅₀ /2):1.7***	0.062*	0.04
Freshwater diatom (N. pelliculosa)	acute	AE C656948	(EC ₅₀ /2): 3.1***	0.062	0.02
Duckweed (L. gibba)	acute	AE C656948	(EC ₅₀ /2): 1.3***	0.062*	0.05
Sheepshead Minnow (C. variegatus)	acute	AE C656948	(LC ₅₀ /10): >0.098**	0.062*	< 0.63
Eastern Oyster (C. virginica)	acute	AE C656948	(LC ₅₀ /2): 0.22***	0.062*	0.28
Saltwater Diatom (S. costatum)	acute	AE C656948	(EC ₅₀ /2): >0.57***	0.062*	< 0.11
Marine amphipods (Leptocheirus	acute	AE C656948	(LC ₅₀ /2): >0.8*** (overlying water)	0.062*	<0.08
plumulosus)	chronic	AE C656948	NOAEC: 0.55 (overlying water)	0.062*	0.11

^{* 80} cm water depth; £15 cm water depth

Table 53 Refined Risk Assessment to Amphibians: Run off (Luna Privilege)

	Exposure	Test Substance	Tox Value For RQ (mg a.i./L)	EEC (mg a.i./L)	RQ
D off	acute	AE C656948	(LC ₅₀ /10): >0.178	0.299*	<1.68
Run off	chronic	AE C656948	NOAEC: 0.135	0.261**	1.93

^{*} peak concentration and ** 21-day EEC in 15 cm water depth

^{**} with an uncertainty factor of 10

^{***} with an uncertainty factor of 2

Table 54 Refined Risk Assessment to Amphibians: Spray Drift (Luna Privilege)

	Airblast early (74% drift)	Airblast late (59% drift)	Ground boom (6% drift)
Application rate (250 g a.i./ha)	185 g a.i./ha	147.5 g a.i./ha	15.00 g a.i/ha
Cumulative application rate (2 applications, 7 d interval)	369.39 g a.i./ha	294.51 g a.i./ha	29.95 g a.i./ha
EEC	0.25 mg a.i./L	0.20 mg a.i./L	0.02mg a.i./L
Acute RQ(LC ₅₀ : 0.178 mg a.i./L)	1.40	1.12	0.11
Chronic RQ (NOEC: 0.135 mg a.i./L)	1.85	1.48	0.15
Risk	yes	yes	no

Table 55 Screening Level and Refined Risk Assessment to Amphibians: Spray Drift from Aerial Application (Potato)

	Screening level (direct overspray)	Aerial application-in off-field (23% drift)
Application rate (400 g a.i./ha)	398.5 g a.i./ha (cumulative)	91.7 g a.i./ha
EEC in 15 cm water depth	0.266 mg a.i./L	0.061 mg a.i./L
Acute RQ(LC ₅₀ : 0.178 mg a.i./L)	1.49	0.34
Chronic RQ (NOEC: 0.135 mg a.i./L)	1.97	0.45
Risk	yes	no

Table 56 Screening Level Risk Assessment to Amphibians (Propulse Fungicide)

Exposure	Test Substance	Tox Value for RQ (mg a.i./L)	EEC (mg a.i./L)	RQ
Acute	AE C656948	(LC ₅₀ /10): 0.178	0.2*	1.12
Chronic	AE C656948	NOAEC: 0.135	0.2*	1.48

^{* 15} cm water depth

Table 57 Refined Risk Assessment to Amphibians: Run Off (Propulse Fungicide)

Exposure	Test Substance	Tox Value For RQ (mg a.i./L)	EEC (mg a.i./L)	RQ
Acute	AE C656948	(LC ₅₀ /10): >0.178	0.179*	<1.00
Chronic	AE C656948	NOAEC: 0.135	0.157**	1.16

^{** 15} cm water depth and 21-day EEC

Table 58 Refined Risk Assessment to Amphibians: Spray Drift (Propulse Fungicide)

Ground Boom (6% drift)
9.00 g a.i/ha
17.97 g a.i./ha
0.002mg a.i./L
0.01
0.01
no

Table 59 Luna Privilege Use (Label) Claims Proposed by Applicant and Whether Acceptable or Unsupported

Proposed use claim	Supported Use
To control powdery mildew on watermelon, apply Luna Privilege at a rate of 150-250 mL/ha at seven to fourteen day intervals.	Supported as proposed
To control botrytis grey mold on watermelon, apply Luna Privilege at a rate of 500 mL/ha at seven to ten day intervals.	Supported as proposed
To control botrytis bunch rot / grey mold on wine grape, apply Luna Privilege at a rate of 500 mL/ha at early bloom and at berry touch to bunch closure.	Supported as proposed
To control white mold on dry bean, apply Luna Privilege at a rate of 300 mL/ha at seven to fourteen day intervals.	Supported as proposed
To control ascochyta blight on dry bean, apply Luna Privilege at a rate of 300 mL/ha at ten to fourteen day intervals.	Supported as proposed
To control mycosphaerella blight on dry bean, apply Luna Privilege at a rate of 300 mL/ha at ten to fourteen day intervals.	Supported as proposed
To control powdery mildew on dry bean, apply Luna Privilege at a rate of 150-250 mL/ha at seven to fourteen day intervals.	Supported as proposed
To control early leaf spot on peanut, apply Luna Privilege at a rate of 250-500 mL/ha at a 14 day intervals.	Supported as proposed
To control late leaf spot on peanut, apply Luna Privilege at a rate of 250-500 mL/ha at a 14 day intervals.	Supported as proposed
To control leaf scab on apple, apply Luna Privilege at a rate of 300 mL/ha at seven to fourteen day intervals.	Supported as proposed
To control early blight on potato, apply Luna Privilege at a rate of 150-300 mL/ha at seven to twelve day intervals.	Supported as proposed
To control powdery mildew on strawberry, apply Luna Privilege at a rate of 500 mL/ha through drip irrigation at five to seven day intervals.	Supported as proposed
To control brown rot blossom blight on sweet and tart cherry, apply Luna Privilege at a rate of 250 mL/ha at fourteen day intervals.	Supported as proposed but limited to three seasonal applications rather than four for resistance management considerations
To control powdery mildew on sweet and tart cherry, apply Luna Privilege at a rate of 150-250 mL/ha at seven to fourteen day intervals.	Supported as proposed but limited to three seasonal applications rather than four for resistance management considerations
To control brown rot blossom blight on tree nuts, apply Luna Privilege at a rate of 250-500 mL/ha at fourteen day intervals.	Supported as proposed on almonds, the only susceptible tree nut in the crop group.

Table 60 Luna Tranquility Fungicide Use (Label) Claims Proposed by Applicant and Whether Acceptable or Unsupported

Proposed use claim	Supported Use
To control powdery mildew on wine grape, apply Luna Tranquility Fungicide at a rate of 600 mL/ha at seven to fourteen day intervals.	Supported as proposed with a limit of four applications per season instead of six.
To control botrytis bunch rot / grey mold on wine grape, Luna Tranquility Fungicide at a rate of 1200 mL/ha at early bloom and at berry touch to bunch closure.	Supported as proposed
To control powdery mildew on apple, apply Luna Tranquility Fungicide at a rate of 600 mL/ha at seven to fourteen day intervals.	Supported as proposed with a limit of four applications per season instead of six.
To control leaf scab on apple, apply Luna Tranquility Fungicide at a rate of 800 mL/ha at seven to fourteen day intervals.	Supported as proposed with a limit of four applications per season instead of six.

Table 61 Propulse Fungicide Use (Label) Claims Proposed by Applicant and Whether Acceptable or Unsupported

Proposed use claim	Supported Use	
To control white mold on dry bean, apply Propulse Fungicide at a rate of 750 mL/ha at seven to fourteen day intervals.	Supported as proposed	
To control ascochyta blight on dry bean, apply Propulse Fungicide at a rate of 500-750 mL/ha at ten to fourteen day intervals.	Supported as proposed	
To control mycosphaerella blight on dry bean, apply Propulse Fungicide at a rate of 500-750 mL/ha at ten to fourteen day intervals.	Supported as proposed	

Table 62 Active Ingredients Currently Registered for Management of Crop Diseases on the Luna Privilege Fungicide, Luna Tranquility Fungicide, and Propulse Fungicide Labels

Crops	Diseases	Active Ingredients (Resistance Management Group)
Watermelon	Powdery mildew ¹	Bacillus subtilis QST 713 (44)
		chlorothalonil (M5)
		difenoconazole (3)
		folpet (M4)
		potassium bicarbonate (NC)
		pyraclostrobin (22)
		Streptomyces lydicus WYEC 108 (NC)
	Botrytis grey mold ¹	ferbam (M3)
		Gliocladium catenulatum J1446 (NC)
		iprodione (2)

Crops	Diseases	Active Ingredients (Resistance Management Group)
Wine Grape	Botrytis bunch rot / Grey	Bacillus subitlis QST 713 (44)
Wille Grape	mold ^{1,3}	boscalid (7) + pyraclostrobin (11)
		fenhexamid (17)
	Powdery mildew ³	Bacillus subtilis QST 713 (44)
	l owdery influew	boscalid (7)
		calcium polysulfide (M2)
		copper oxychloride (M2)
		difenoconazole (3)
		folpet (M4)
		kresoxim-methyl (11)
		metrafenone (U8)
		myclobutanil (3)
		potassium bicarbonate (NC)
		pyraclostrobin (11) + boscalid (7)
		quinoxyfen (13)
		sulphur (M2)
D D	11.2	trifloxystrobin (11)
Dry Bean	White mold ^{1,2}	Bacillus subtilis QST 713 (44)
(Including Chickpea		boscalid (7)
and Lentil)		Coniothyrium minitans CON/M/91-08 (NC)
		cyprodinil (9) + fludioxonil (12)
		dicloran (14)
		fluazinam (29)
		iprodione (2)
	13	vinclozolin (2)
	Ascochyta blight ^{1,2}	azoxystrobin (11)
	12	pyraclostrobin (11)
	Mycosphaerella blight ^{1,2}	azoxystrobin (11)
	Powdery mildew ¹	pyraclostrobin (11)
		azoxystrobin (11) + propiconazole (3)
		propiconazole (3)
		pyraclostrobin (11)
Peanut	Early Leaf Spot ¹	Bacillus subtilis QST 713 (44)
		prothioconazole (3)
	Late Leaf Spot ¹	Bacillus subtilis QST 713 (44)
Apple	Leaf scab ^{1,3}	Bacillus subtilis QST 713 (44)
		boscalid (7) + pyraclostrogbin (11)
		calcium polysulphide (M2)
		captan (M4)
		cyprodinil (9)
		difenoconazole (3)
		dodine (M7)
		ferbam (M3)
		fluazinam (29)

Crops	Diseases	Active Ingredients	
		(Resistance Management Group)	
		flusilazole (3)	
		folpet (M4)	
		kresoxim-methyl (11)	
		mancozeb (M3)	
		mancozeb (M3) + myclobutanil (3)	
		metiram (M2)	
		myclobutanil (3)	
		pyrimethanil (9)	
		penthiopyrad (7)	
		sulphur (M2)	
		thiophanate-methyl (1)	
		thiram (M3)	
		trifloxystrobin (11)	
		ziram (M3)	
	Powdery mildew ³	Bacillus subtilis QST 713 (44)	
		boscalid (7) + pyraclostrogbin (11)	
		calcium polysulphide (M2)	
		chlorothalonil (M5)	
		cyprodinil (9)	
		difenoconazole (3)	
		flusilazole (3)	
		kresoxim-methyl (11)	
		myclobutanil (3)	
		sulphur (M2)	
		thiophanate-methyl (1)	
		trifloxystrobin (11)	
Potato	Early blight ¹	azoxystrobin (11)	
Otato	Carry bright	Bacillus subtilis QST 713 (44)	
		boscalid (7)	
		captan (M4)	
		chlorothalonil (M5)	
		copper – different salts (M1)	
		difenoconazole (3)	
		dimethomorph (40) + mancozeb (M3)	
		famoxadone (11) + cymoxanil (27)	
		mancozeb (M3)	
		mancozeb (M3) + zoxamide (22)	
		maneb (M3)	
		metalaxyl (4) + chlorothalonil (M5)	
		metalaxyl (4) + mancozeb (M3)	
		metiram (M3)	
		pyraclostrobin (11)	
		zineb (M3)	
		zoxamide (22)	

Crops	Diseases	Active Ingredients (Resistance Management Group)
Strawberry	Powdery mildew ¹	boscalid (7) + pyraclostrobin (11)
		calcium polysulphide (M2)
		citric acid (NC) + lactic acid (NC)
		myclobutanil (3)
		quinoxyfen (13)
		Streptomyces lydicus WYEC 108 (NC)
Cherry	Brown rot blossom blight	Bacillus subtilis QST 713 (44)
		boscalid (7)
		boscalid (7) + pyraclostrobin (11)
		chlorothalonil (M5)
		cyprodinil (9)
		dicloran (14)
		fenhexamid (17)
		pyraclostrobin (11)
	Powdery mildew ¹	boscalid (7) + pyraclostrobin (11)
		quinoxyfen (13)
Almond	Brown rot / blossom blight ¹	chlorothalonil ⁴ (M5)

claim appears on the Luna Privilege label

Table 63 TSMP considerations-comparison to TSMP Track 1 criteria

TSMP Track 1 Criteria TSMP Tra		rack 1 Criterion value	Active Ingredient Endpoints	
Toxic or toxic equivalent as defined by the <i>Canadian</i> Environmental Protection Act ¹	by the Canadian			
Predominantly anthropogenic ²	yes			
Persistence ³ :	soil	half-life: ≥ 182 days	field DT ₅₀ : 539 days	
	water	half-life: ≥ 182 days	half-life: 1470 days (water+sediment)	
	sediment	half-life: ≥ 365 days	not available	
	air	half-life ≥ 2 days or evidence of long range transport	1.7 to 2.6 days	
Bioaccumulation ⁴	log K _{OW} ≥		3.3	
	BCF ≥ 5000		18	
	$BAF \ge 5000$			
Is the chemical a TSMP Track 1 met)?	substance (a	all four criteria must be	no, does not meet TSMP Track 1 criteria.	

All pesticides will be considered toxic or toxic equivalent for the purpose of initially assessing a pesticide against the TSMP criteria. Assessment of the toxicity criterion may be refined if required (that is, all other TSMP criteria are met).

²claim appears on the Propulse Fungicide label

³claim appears on the Luna Tranquility Fungicide label

⁴ registered for ornamental applications only

² The policy considers a substance "predominantly anthropogenic" if, based on expert judgement, its concentration

- in the environment medium is largely due to human activity, rather than to natural sources or releases.
- If the pesticide and/or the transformation product(s) meet one persistence criterion identified for one media (soil, water, sediment or air) than the criterion for persistence is considered to be met.
- Field data (for example, BAFs) are preferred over laboratory data (for example, BCFs) which, in turn, are preferred over chemical properties (for example, logK_{ow}).

Appendix II Supplemental Maximum Residue Limit Information— International Situation and Trade Implications

Fluopyram is a new active ingredient, which is concurrently being registered in the United States. American tolerances (40 CFR Part 180) and Codex MRLs established for fluopyram differ from the Canadian maximum residue limits (MRLs) as shown in table below.

Table 1 Differences Between MRLs in Canada and in Other Jurisdictions

Commodity Commodity	Canada (ppm)	U.S. (ppm)	Codex* (ppm)
Wine grapes	2.0	2.0	2 (Grapes); 5 (Dried grapes)
Canola	1.8	1.8	
Crop Group 15 (except rice) – Cereal Grains, except rice; Strawberries	1.5	1.5	
Cherries	1.5	0.6	
Bananas; Watermelon	1.0	1.0	
Dry chickpeas and dry lentils	0.4	None	
Apples	0.3	0.3	
Sugar beet roots	0.1	0.04	
Dry soybeans	0.1	0.1	
Grain lupin, dry kidney beans, dry lima beans, dry navy beans, dry pink beans, dry pinto beans, dry tepary beans, dry beans, dry adzuki beans, dry blackeyed peas, dry catjang seed, dry cowpea seed, dry moth beans, dry mung beans, dry rice beans, dry southern peas, dry urd beans, dry broad beans, dry guar seed, dry lablab beans	0.09	0.09	Not reviewed by Codex
Crop Group 14 – Tree Nuts Group	0.05	0.05	
Crop Subgroup 1C – Tuberous and Corm Vegetables Subgroup	0.02	0.02 (potato)	
Peanuts	0.02	0.02	
Undelinted cotton seeds	0.01	0.01	
Eggs	0.06	0.25	
Meat byproducts of poultry	0.10	0.60	
Fat of poultry	0.03	0.20	
Meat of poultry	0.03	0.15	
Milk	0.06	0.07	0.07
Meat byproducts of cattle, goats, horses and sheep	0.40	1.1	0.7 (Edible
Fat of cattle, goats, horses and sheep	0.05	0.11	offal,
Meat of cattle, goats, horses and sheep	0.05	0.15	mammalian);
Meat byproducts of hogs	0.03	0.70	0.1 (Meat from
Fat and meat of hogs	0.02	0.05	mammals other than marine mammals)

^{*} Codex is an international organization under the auspices of the United Nations that develops international food standards, including MRLs.

MRLs may vary from one country to another for a number of reasons, including differences in pesticide use patterns and the locations of the field crop trials used to generate residue chemistry data. For animal commodities, differences in MRLs can be due to different livestock feed items and practices.

Under the North American Free Trade Agreement NAFTA, Canada, the United States and Mexico are committed to resolving MRL discrepancies to the broadest extent possible. Harmonization will standardize the protection of human health across North America and promote the free trade of safe food products. Until harmonization is achieved, the Canadian MRLs specified in this document are necessary. The differences in MRLs outlined above are not expected to impact businesses negatively or adversely affect international competitiveness of Canadian firms or to negatively affect any regions of Canada.

References

A. List of Studies/Information Submitted by Registrant

1. Chemistry

PMRA Document	Reference
Number 1599498	2008, 1st amendment to report no. 20050612.01 - AE C656948; Substance, pure - AE C656948 00 1B99 0001- Vapour pressure A.4. (OECD 104), DACO: 2.14.9,IIA 2.3.1
1599499	2008, 1st amendment to report no. 20070286.01 - Fluopyram, AE C656948 - Melting point A.1. (OECD 102), Boiling point A.2. (OECD 103), Thermal stability (OECD 113), DACO: 2.14.13,2.14.4,2.14.5,IIA 2.1.1,IIA 2.1.2,IIA 2.1.3
1599500	2008, 1st amendment to report no. 20070286.03 - Fluopyram, AE C656948 - Explosive properties A.14., DACO: 2.16,IIA 2.13
1599502	2008, 1st amendment to report no. 20070359.01 - Fluopyram, AE C656948, pure substance - Product code: AE C656948 001B99 0001 - Melting point A.1. (OECD 102), Boiling point A.2. (OECD 103), Thermal stability (OECD 113), DACO: 2.14.13,2.14.4,2.14.5,IIA
1599504	2008, 1st amendment to storage stability of AE C656948, DACO: 2.14.14,IIA 2.17.1
1599549	2008, AE C656948 Determination of active substance in technical material HPLC - internal standard, DACO: 2.13.1,IIA 4.2.1
1599565	2008, AE C656948 - By-products in technical grade active substance [CBI removed], DACO: 2.13.4,IIA 4.2.3 CBI
1599628	2008, Analytical procedure for the Karl Fischer water determination, DACO: 2.13.4,IIA 4.2.3 CBI
1599637	2007, Composition statement - Technical grade active substance - Fluopyram - Fluopyram TC, AE C656948, DACO: 2.12.2,2.3.1,2.4,2.5,2.6,2.7,2.8,2.9,IIA 1.10.1 CBI
1599641	2008, Determination of [CBI removed] in technical grade active substance [C BI removed], DACO: 2.13.4,IIA 4.2.3 CBI
1599643	2008, Determination of the pH-value of AE C656948 pure substance, DACO: 2.16,IIA 2.16
1599675	2008, Determination of the storage stability of AE C656948 and its metabolites [CBI removed] in soil - results for an interval of 0 to 323 days, DACO: 2
1599676	2008, Determination of [CBI removed] in active ingredient of agrochemicals [CBI removed], DACO: 2.13.4,IIA 4.2.3 CBI
1599678	2008, Dissociation constant of fluopyram (AE C656948) in water, DACO: 2.14.10,8.2.3.2,IIA 2.9.5
1599742	2008, Fluopyram (AE C656948) - Statement on the ecotoxicological assessment of the proposed technical specification, DACO: 2.13.3,IIA 1.11.2 CBI
1599743	2008, Fluopyram (AE C656948) - Toxicological equivalence assessment of the technical specification with the material tested in toxicity studies, DACO: 2.13.3,IIA 1.11.2 CBI

1599744	2008, Fluopyram (AE C656948): Statement on the dielectric breakdown voltage according to OPPTS 830.6321, DACO: 2.16,IIA 2.18
1599745	2008, Fluopyram (AE C656948): Statement on the miscibility according to OPPTS 830.6319, DACO: 2.16,IIA 2.18
1599746	2008, Fluopyram (AE C656948): Statement on the pH independence of the partition coefficient 1-octanol / water, DACO: 2.14.11,IIA 2.8.2
1599747	2008, Fluopyram (AE C656948): Statement on the viscosity according to OPPTS 830.7100, DACO: 2.16,IIA 2.18
1599748	2008, Fluopyram (AEC 656948) - Technical grade active substance - Discussion of the formation of impurities, DACO: 2.11.1,2.11.3,2.11.4,2.12.2,IIA 1.10.1,IIA 1.8.1 CBI
1599749	2008, Fluopyram (AEC 656948) - Technical grade active substance - Manufacturing process, DACO: 2.11.1,2.11.2,2.11.3,2.11.4,IIA 1.8.1,IIA 1.8.2 CBI
1599755	2008, Fluopyram, AE C656948 - Auto-flammability (Bowes-Cameron-Cage test), DACO: 2.16,IIA 2.11.2
1599756	2008, Fluopyram, AE C656948 - Auto-flammability (solids - determination of relative self-ignition temperature) A.16., DACO: 2.16,IIA 2.11.2
1599757	2008, Fluopyram, AE C656948 - Flammability (solids) A.10., DACO: 2.16,IIA 2.11.1
1599758	2008, Fluopyram, AE C656948 - Oxidizing properties A.17., DACO: 2.16,IIA 2.13
1599764	2008, Henry's law constant of fluopyram (AE C656948) at pH 4, pH 7 and pH 9, DACO: 2.16,IIA 2.3.2
1599765	2008, Henry's law constant of fluopyram (AE C656948), DACO: 2.16,IIA 2.3.2
1599776	2008, Material accountability chronic tox sample of fluopyram, DACO: 2.13.3,IIA 1.11.2 CBI
1599777	2008, Material accountability of Fluopyram batches used in toxicological and ecotoxicological studies, DACO: 2.13.3,IIA 1.11.2 CBI
1599778	2008, Material accountability of fluopyram manufactured at Dormagen / Germany, DACO: 2.13.3,IIA 1.11.1 CBI
1599797	2008, Particle size, fiber length and diameter distribution of fluopyram technical, DACO: 2.16,IIA 2.18
1599798	2008, Partition coefficients 1-octanol / water of AE C656948 - (shake flask method), DACO: 2.14.11,IIA 2.8.1
1599807	2008, Physical characteristics color, physical state and odor of fluopyram (AE C656948) Technical substance and pure substance, DACO: 2.14.1,2.14.2,2.14.3,IIA 2.4.1,IIA 2.4.2
1599810	2008, Relative density of fluopyram (AE C656948), pure substance and technical substance, DACO: 2.14.6,IIA 2.2
1599814	2008, Solubility in organic solvents AE C656948 pure substance, DACO: 2.14.8,IIA 2.7
1599816	2007, Spectral data set of AE C656948, DACO: 2.13.2,2.14.12,IIA 2.5.1.1,IIA 2.5.1.2,IIA 2.5.1.3,IIA 2.5.1.4,IIA 2.5.1.5
1599817	2008, Stability to normal and elevated temperature, metals, and metal ions of Fluopyram, DACO: 2.14.13,IIA 2.17.2
1599820	2008, Storage stability of AE C656948 - Amendment no.1, DACO: 2.14.14,IIA 2.17.1

1599822	2008, Surface tension of fluopyram (AE C656948) technical substance, DACO: 2.16,IIA 2.14
1599833	2008, The oxidation or reduction properties of fluopyram (AE C656948), technical substance, DACO: 2.16,IIA 2.18
1599869	2008, UV/VIS spectral data set of fluopyram, DACO: 2.13.2,2.14.12,IIA 2.5.1.1,IIA 2.5.1.5
1599872	2008, Validation of [CBI removed] - Fluopyram - Byproducts in technical grade active substance [CBI removed], DACO: 2.13.4,IIA 4.2.3 CBI
1599873	2008, Validation of HPLC-method AM002705MP1 AE C656948 assay of technical grade active substance, DACO: 2.13.1,IIA 4.2.1
1599874	2008, Validation of [CBI removed] - Fluopyram - Byproducts in technical grade active substance [CBI removed], DACO: 2.13.4,IIA 4.2.3 CBI
1599875	2008, Validation of [CBI removed] - determination of [CBI removed] in technical grade active [CBI removed] in AE C656948 [CBI removed], DACO: 2.13.4,IIA 4.2.3 CBI
1599876	2008, Water solubility of AE C656948 at pH 4, pH 7, pH 9 and in distilled water (Flask method), DACO: 2.14.7,IIA 2.6
1745793	Draft Assessment Report of Fluopyram, DACO: 2.0,3.0 CBI
1762436	2009, Position Paper - Addressing the Identity of the Impurity [CBI removed], DACO: 2.13.2 CBI
1908576	2010, Composition Statement - Technical grade active substance - Fluopyram TC, AE C656948, DACO: 2.12.2,2.13.4,IIA 1.10.1,IIA 1.10.2 CBI
1908577	2010, Fluopyram (AEC 656948) Technical grade active substance - Discussion of the formation of impurities, DACO: 2.12.2,2.13.4,IIA 1.10.2 CBI
1908579	2010, 2nd Amendment to material accountability of fluopyram manufactured at Dormagen / Germany - Five batches of technical fluopyram, DACO: 2.13.3,IIA 1.11.1 CBI
1908580	2010, Material accountability of Fluopyram - re-analysis of impurities in 5 technical batches, DACO: 2.13.3,IIA 1.11.1 CBI
1908583	2010, 1st amendment to material accountability of chronic tox sample of fluopyram, DACO: 2.13.3,IIA 1.11.2 CBI
1908588	2010, 1st amendment to material accountability of fluopyram batches used in toxicological and ecotoxicological studies, DACO: 2.13.3,IIA 1.11.2 CBI
1908589	2010, Fluopyram (AE C656948) - Toxicological equivalence assessment of the technical specification with the material tested in toxicity studies - 1st amendment, DACO: 2.13.3,IIA 1.11.2 CBI
1908590	2010, Fluopyram (AE C656948) - Statement on the ecotoxicological assessment of the proposed technical specification, DACO: 2.13.3,IIA 1.11.2
1908593	2010, Material accountability of fluopyram (AE C656948) - Re-analysis of impurities in chronic tox sample, DACO: 2.13.3,IIA 1.11.2 CBI
1908594	2010, Material accountability of fluopyram batches used in toxicological and ecotoxicological studies - Re-analysis of impurities, DACO: 2.13.3,IIA 1.11.2 CBI
1908596	2010, Analytical Method, AE C656948 Impurities in technical grade substance [CBI removed], DACO: 2.13.4,IIA 4.2.3 CBI
1908597	2010, Validation of [CBI removed] Impurities on technical grade active substance [CBI removed], DACO: 2.13.4,IIA 4.2.3 CBI
1908598	2010, Clarification regarding impurities in the material accountability studies of fluopyram, DACO: 2.13.4,IIA 4.2.3 CBI

1599620 2006, Analytical method 00973 for the determination of residues of AE C656948 in soil by HPLC-MS/MS, DACO: 8.2.2.1, IIA 4.4 2007, Analytical method 01023 for the determination of residues of AE C656948 1599622 and its metabolites [CBI removed] in soil by HPLC-MS/MS, DACO: 8.2.2.1, IIA 4.4 1599623 2007, Analytical method 01051 for the determination of fluopyram (AE C656948) in drinking and surface water by HPLC-MS/MS, DACO: 8.2.2.3, IIA 4.5 2008, Analytical Method 01068 for the determination of residues of AE C656948 1599625 in soil by HPLC-MS/MS, DACO: 8.2.2.1, IIA 4.4 2008, Analytical method for the determination of residues of AE C656948 and its 1599627 metabolites AE C656948-benzamide, AE C656948-7-hydroxy, and AE C656948-PCA in soil and sediment using LC/MS/MS, DACO: 8.2.2.2, IIA 4.6 2008, Determination of fluopyram (AE C656948) in water by LC/MS/MS, DACO: 1599642 8.2.2.3,IIA 4.5 1599766 2008. Independent laboratory validation of analytical method 01023 for the determination of residues of AE C656948 and its metabolites AE C656948benzamide (AE F148815), AE C656948-7-hydroxy (BCS-AA-10065) and AE C656948-PCA in soil by HPLC-MS/MS on soil 2008. Independent laboratory validation of analytical method 01051 for the 1599767 determination of fluopyram (AE C656948) in drinking and surface water by HPLC-MS/MS, DACO: 8.2.2.3.IIA 4.5 1599279 2008, Composition statement - Plant protection product - Fluopyram SC 500 (500 g/L), DACO: 3.3.2, IIIA 1.4.1 CBI 2008. Product chemistry of fluopyram 500 SC, DACO: 3.2.1,3.3.1,3.3.2,IIIA 1.4.2 1599280 CBI 1599283 2008, Manufacturing process description for fluopyram 500 SC, DACO: 3.2.2,IIIA 1.4.5.1 CBI 2008, Physical, chemical and technical properties of fluopyram SC 500 (500 g/L), 1599304 DACO: 3.5.1,3.5.2,3.5.3,3.5.6,3.5.7,3.5.9,3.7,8.2.2.1,8.2.3.6,IIIA 2.1,IIIA 2.4.2,IIIA 2.5.1,IIIA 2.5.2,IIIA 2.5.3,IIIA 2.6.1,IIIA 2.8.2,IIIA 2.8.3.1,IIIA 2.8.3.2.IIIA 2.8.5. 1599305 2008, Miscibility of fluopyram 500 SC, DACO: 3.5.13, IIIA 2.11 2008, Dielectric breakdown voltage of fluopyram 500 SC, DACO: 3.5.15,IIIA 2.12 1599306 1599307 2007, Storage stability of fluopyram SC 500 (500 g/L) - [Packaging material: HDPE] - Interim report (8 weeks), DACO: 3.5.10,3.5.14,IIIA 2.13,IIIA 2.7.1,IIIA 2.7.2, IIIA 2.7.3, IIIA 2.7.4, IIIA 2.7.5 2008, Container material of fluopyram SC 500, DACO: 3.5.5,IIIA 2.14 1599308 1599309 2008, Safety relevant technical properties of fluopyram SC 500 g/L -Final report-, DACO: 3.5.11,3.5.12,3.5.8,IIIA 2.2.1,IIIA 2.2.2,IIIA 2.3.1,IIIA 2.3.3 2006, Determination of AE C656948 in formulations - Assay - GLC, internal 1599312 standard, DACO: 3.4.1, IIIA 5.2.1 2008, Validation of GLC-method AM005005MF1 -determination of AE C656948 1599313 in formulations-, DACO: 3.4.1, IIIA 5.2.1 2008, Validation of GLC-method AM005005MF1 - Determination of AE C656948 1599314 in formulations, DACO: 3.4.1, IIIA 5.2.1 2009, Discussion of the Formation of impurities of Fluopyram 500 SC, DACO: 1764319 3.2.3 CBI

1983757	2009, Storage stability data of fluopyram SC 500 (500 g/L) - packaging material: HDPE, DACO: 3.5.10,3.5.14 CBI
1670065	2008, Composition statement - Plant protection product - Fluopyram + pyrimethanil SC 500 (125 + 375 g/L), DACO: 3.3.2,IIIA 1.4.1 CBI
1670068	2008, Product chemistry of fluopyram + pyrimethanil 500 SC, DACO: 3.2.2,IIIA 1.4.5.1 CBI
1670069	2008, Discussion of the formation of impurities of fluopyram & pyrimethanil - SC 500 (125 + 375 g/L), DACO: 3.2.3,IIIA 1.4.5.2 CBI
1670071	2008, Physical and chemical properties of fluopyram + pyrimethanil 500 SC, DACO: 3.5.1,3.5.2,3.5.3,3.5.6,3.5.7,3.5.9,IIIA 2.1,IIIA 2.4.2,IIIA 2.5.2,IIIA 2.6.1
1670072	2008, Miscibility of fluopyram & pyrimethanil - SC 500 (125+ 375 g/L), DACO: 3.5.13, IIIA 2.11
1670073	2008, Dielectric breakdown voltage of fluopyram & pyrimethanil - SC 500 (125+375 g/L), DACO: 3.5.15,IIIA 2.12
1670074	2008, Container material of fluopyram & pyrimethanil - SC 500 (125+ 375 g/L), DACO: 3.5.5,IIIA 2.14
1670075	2008, Determination of safety-relevant data of fluopyram + pyrimethanil SC 500 (125+375 g/L), DACO: 3.5.11,3.5.12,3.5.8,IIIA 2.2.1,IIIA 2.2.2,IIIA 2.3.1,IIIA 2.3.3
1670078	2008, Determination of fluopyram and pyrimethanil in formulations - assay - GLC internal standard, DACO: 3.4.1,IIIA 5.2.1,IIIA 5.2.2
1670079	2008, Validation of GLC-method AM010007MF1 - determination of fluopyram and pyrimethanil in formulations, DACO: 3.4.1,IIIA 5.2.1,IIIA 5.2.2
1838560	2009, Determination of pH of Water, Flowables, and Aqueous Solutions, DACO: 3.5.7 CBI
1838561	2008, Brookfield Viscosity, DACO: 3.5.9 CBI
1838565	2009, Storage Stability Data of fluopyram + pyrimethanil SC 500 (125+375 g/L), DACO: 3.5.10,3.5.14 CBI
1670779	2008, Product chemistry of fluopyram + prothioconazole 400 SC, DACO: 3.2.2,3.3.1,3.3.2,3.4.1,3.5.1,3.5.10,3.5.11,3.5.12,3.5.13,3.5.15,3.5.2,3.5.3,3.5.6,3.5 7,3.5.8,3.5.9,IIIA 1.4.1,IIIA 1.4.2,IIIA 1.4.5.1,IIIA 2.1,IIIA 2.11,IIIA 2.12,IIIA 2.2.1,IIIA 2.2.2
1670784	2008, Container material of fluopyram & prothioconazole - SC 400 (200 + 200 g/L), DACO: 3.5.5,IIIA 2.14
1838577	2009, Determination of pH of Water, Flowables, and Aqueous Solutions, DACO: 3.5.7 CBI
1838578	2008, Brookfield Viscosity, DACO: 3.5.9 CBI
2174226	2012, Storage Stability and Corrosion Characteristics of Fluopyram + Prothioconazole SC 400 (Propulse 400 SC), DACO: 3.5.10,3.5.14 CBI

2.0 Human and Animal Health

PMRA Document Number	Reference
1599505	2003, 28-day toxicity study in the rat by dietary administration Code: AE 1344122, DACO: 4.8,IIA 5.8
1599513	2008, [Phenyl-UL-14C]AE 656948: Absorption, distribution, excretion and metabolism in the rat, DACO: 4.5.9,IIA 5.1.1
1599517	2008, [Phenyl-UL-14C]AE C656948: Distribution of the total radioactivity in male and female rats determined by quantitative whole body autoradiography (QWBA), determination of the exhaled 14CO2 and metabolic profiling in excreta, DACO: 4.5.9,IIA 5.1.1
1599524	2008, [Pyridyl-2,6-14C]AE C656948 - Metabolism in organs and tissues of male and female rats (three timepoints), DACO: 4.5.9,IIA 5.1.1
1599526	2008, [Pyridyl-2,6-14C]AE C656948: Absorption, distribution, excretion and metabolism in the rat, DACO: 4.5.9,IIA 5.1.1
1599529	2008, [Pyridyl-2,6-14C]AE C656948: Distribution of the total radioactivity in male and female rats determined by quantitative whole body autoradiography (QWBA), determination of the exhaled 14CO2, DACO: 4.5.9,IIA 5.1.1
1599533	2007, A subacute dermal toxicity study in rats with technical grade AE C656948, DACO: 4.3.5,IIA 5.3.7
1599534	2008, A subchronic neurotoxicity screening study with technical grade AE C656948 in Wistar rats, DACO: 4.5.13,IIA 5.7.4
1599545	2003, Acute toxicity in the rat after oral administration AE 1344122 Project AE C638206, DACO: 4.8, IIA 5.8
1599547	2003, AE 1344122 (metabolite of AE C638206): Induction of chromosome aberrations in cultured human peripheral blood Imphocytes, DACO: 4.8,IIA 5.8
1599548	2008, AE C656948 - Chronic toxicity study in the dog by dietary administration, DACO: 4.3.2,IIA 5.3.4
1599551	2008, AE C656948 (fluopyram) - In vitro studies on the potential interactions with thyroid peroxidase-catalyzed reactions, DACO: 4.8,IIA 5.5.4
1599552	2007, AE C656948 (project: AE C656948) - In vitro chromosome aberration test with Chinese hamster V79 cells, DACO: 4.5.6,IIA 5.4.2
1599553	2008, AE C656948 (project: fluopyram) - Salmonella/microsome test - Plate incorporation and preincubation method, DACO: 4.5.4,IIA 5.4.1
1599555	2008, AE C656948 - 90-day toxicity study in the dog by dietary administration, DACO: 4.3.2,IIA 5.3.3
1599556	2008, AE C656948 - 90-day toxicity study in the mouse by dietary administration, DACO: 4.3.1,IIA 5.3.2
1599557	2007, AE C656948 - 90-day toxicity study in the rat by dietary administration, DACO: 4.3.1,IIA 5.3.2
1599558	2007, AE C656948 - Acute eye irritation on rabbits, DACO: 4.2.4,IIA 5.2.5
1599559	2007, AE C656948 - Acute inhalation toxicity in rats, DACO: 4.2.3,IIA 5.2.3
1599561	2007, AE C656948 - Acute skin irritation/corrosion on rabbits, DACO: 4.2.5,IIA 5.2.4

1599563	2007, AE C656948 - Acute toxicity in the rat after dermal application, DACO: 4.2.2,IIA 5.2.2
1599564	2007, AE C656948 - Acute toxicity in the rat after oral administration, DACO: 4.2.1,IIA 5.2.1
1599571	2008, AE C656948 - Developmental toxicity study in the rabbit by gavage, DACO: 4.5.3,IIA 5.6.11
1599573	2008, AE C656948 - Evaluation of potential dermal sensitization in the local lymph node assay in the mouse, DACO: 4.2.6,IIA 5.2.6
1599574	2008, AE C656948 - Exploratory 28-day toxicity study in the rat by dietary administration, DACO: 4.3.3,IIA 5.3.1
1599576	2008, AE C656948 - Mechanistic 14-day toxicity study in the mouse by dietary administration (hepatotoxicity and thyroid hormone investigations), DACO: 4.8,IIA 5.5.4
1599577	2007, AE C656948 - Micronucleus-test on the male mouse, DACO: 4.5.7,IIA 5.4.4
1599578	2008, AE C656948 - Preliminary 28-day toxicity study in the dog by gavage, DACO: 4.3.3,IIA 5.3.1
1599579	2008, AE C656948 - Preliminary 28-day toxicity study in the mouse by dietary administration, DACO: 4.3.3,IIA 5.3.1
1599580	2007, AE C656948 - Salmonela/microsome test plate incorporation and preincubation method, DACO: 4.5.4,IIA 5.4.1
1599581	2007, AE C656948 - V79/HPRT-test in vitro the detection of induced forward mutations, DACO: 4.5.5,IIA 5.4.3
1599610	2008, AE C656948: Developmental toxicity study in the rat by gavage, DACO: 4.5.2,IIA 5.6.10
1599611	2003, AE C657188 (metabolite of AE C638206): Induction of chromosome aberrations in cultured human peripheral blood lymphocytes, DACO: 4.8,IIA 5.8
1599612	2008, AE C657188 (PCA) Preliminary 28-day toxicity study in the rat by dietary administration Version 2, DACO: 4.8,IIA 5.8
1599613	2003, AE C657188 - V79/HPRT-test in vitro for the detection of induced forward mutations, DACO: 4.8,IIA 5.8
1599617	1984, Chen, H. J., Age and sex difference in serum and pituitary thyrotropin concentrations in the rat: influence by pituitary adenoma, Experimental Gerontology, Vol. 19, pp. 1-6, DACO: 4.8,IIA 5.5.4
1599618	2007, An acute oral neurotoxicity screening study with technical grade AE C656948 in Wistar rats, DACO: 4.5.12,IIA 5.7.1
1599630	2000, Bacterial mutation assay AE C657188 (plant metabolite of AE C638206) Code: AE C657188 00 1B99 0002, DACO: 4.8,IIA 5.8
1599631	2008, Boring, C. C.; Squires, T. S.; Tong, T.; Montgomery, S., Cancer statistics, 1994, A Cancer Journal for Clinicians, 44, pp. 7-26, DACO: 4.8,IIA 5.5.4
1599632	2008, Carcinogenicity study of AE C656948 in the C57BL/6J mouse by dietary administration, DACO: 4.4.3,IIA 5.5.3
1599635	2008, Chronic toxicity and carcinogenicity study of AE C656948 in the Wistar rat by dietary administration, DACO: 4.4.2,4.4.4,IIA 5.5.2
1599741	2008, Fluopyram (AE C656948) - 7-day mechanistic study in the female Wistar rat by dietary administration, DACO: 4.8,IIA 5.5.4

2006, Holsapple, M.; Pitot, H. C.; Cohen, S. H.; Boobis, A. R.; Klanig, J. E.; 1599762 Pastoor, T.; Dellarco, V. L.; Dragan, Y. P., Forum - Mode of action in relevance of rodent liver tumors to human cancer risk, Toxicological Sciences 89(1), pp. 51-56, DACO: 4.8, IIA 5.5.4 2003, Moore, J. T.; Moore, L. B.; Maglich, J. M.; Kliewer, S. A., Functional and 1599763 structural comparison of PXR and CAR, Biochimica et Ciophysics Acta 1619, pp. 235-238, DACO: 4.8, IIA 5.5.4 1599799 1992, Capen, C. C., Pathophysiology of chemical injury of the thyroid gland, Toxicology Letters, 64/65, pp. 381-388. DACO: 4.8,IIA 5.5.4 1599800 2000, Kelly, G., Peripheral metabolism of thyroid hormones: A Review, Alternative Medicine Review, 5(4), pp. 306-333, DACO: 4.8, IIA 5.5.4 1599802 2008, Phenobarbital - 7-day mechanistic study in the female Wistar rat by gavage, DACO: 4.8, IIA 5.5.4 1599803 2008, Phenobarbital - Mechanistic 14-day toxicity study in the mouse by oral gavage (hepatotoxicity and thyroid hormone investigations), DACO: 4.8, IIA 5.5.4 1599804 1996, Whysner, J.; Ross, P. M.; Williams, G. M., Phenobarbital mechanistic data and risk assessment: Enzyme induction, enhanced cell proliferation, and tumor promotion, Pharmacol. Ther., 71(1/2), pp. 153-191, DACO: 4.8, IIA 5.5.4 2000, Rat acute oral toxicity AE C657188 (plant metabolite of AE C638206) 1599809 Code: AE C657188 00 1B99 0002, DACO: 4.8, IIA 5.8 1599812 1998, Hill, R. N.; Crisp, T. M.; Hurley, P. M.; Rosenthal, S. L.; Singh, D. V., Risk assessment of thyroid follicular cell tumors. Environmental Health Perspectives. 106(8), DACO: 4.8, IIA 5.5.4 1599813 2003, Salmonella/microsome test - Plate incorporation and preincubation method Code: AE 1344122, DACO: 4.8, IIA 5.8 1599815 2001, Anon., Some thyrotropic agents - Summary of data reported and evaluation, DACO: 4.8, IIA 5.5.4 1599823 2008, Technical grade AE C656948: A dose range-finding reproductive toxicity study in the Wistar rat, DACO: 4.5.1, IIA 5.6.1 2008, Technical grade AE C656948: A two generation reproductive toxicity study 1599824 in the Wistar rat, DACO: 4.5.1, IIA 5.6.1 1599826 2008, Technical grade AE C656948: A two generation reproductive toxicity study in the Wistar rat, DACO: 4.5.1, IIA 5.6.1 1599827 2008, Technical grade AE C656948: A two generation reproductive toxicity study in the Wistar rat, DACO: 4.5.1, IIA 5.6.1 2008, Technical grade AE C656948: A two generation reproductive toxicity study 1599828 in the Wistar rat, DACO: 4.5.1, IIA 5.6.1 1599829 2008, Technical grade AE C656948: A two generation reproductive toxicity study in the Wistar rat, DACO: 4.5.1, IIA 5.6.1 1599830 2008, Technical grade AE C656948: A two generation reproductive toxicity study in the Wistar rat, DACO: 4.5.1, IIA 5.6.1 1599831 1991, Curran, P. G.; DeGroot, L. J., The effect of hepatic enzyme-inducing drugs on thyroid hormones and the thyroid gland, Endocrine Reviews, 12(2), pp. 135-150, DACO: 4.8, IIA 5.5.4 1599832 1989, McClain, R. M.; Levin, A. A.; Posch, R.; Downing, J. C., The effect of phenobarbital on the metabolism and excretion of Thyroxine in rats, Toxicology

and Applied Pharmacology, 99, pp. 216-228, DACO: 4.8, IIA 5.5.4

1599870	2003, V79/HPRT-test in vitro for the detection of induced forward mutations Code: AE 1344122 (metabolite of AE C628206), DACO: 4.8, IIA 5.8
1654271	1973, Bastomsky, C. H., The biliary excretion of thyroxine and its glucuronic acid conjugate in normal and gunn rats, Endo. 92(1), pp. 35-40, DACO: 4.8,IIA 5.5.4
1654272	2008, AE C656948 - Mechanistic 3-day toxicity study in the male mouse (pharmacokinetic investigations of the clearance of intravenously administered
1654273	125I-thyroxine), DACO: 4.8,IIA 5.5.4 2008, AE C656948 - Mechanistic 3-day toxicity study in the male mouse (QPCR investigations of gene transcripts in the liver), DACO: 4.8,IIA 5.5.4
1661145	2008, AE C656948 - Mechanistic 3-day toxicity study in the male mouse (pharmacokinetic investigations of the clearance of intravenously administered 1251-thyroxine), DACO: 4.8,IIA 5.5.4
1661146	2008, AE C656948 - Mechanistic 3-day toxicity study in the male mouse (QPCR investigations of gene transcripts in the liver), DACO: 4.8,IIA 5.5.4
1729272	2009, AE C656948 Definitive Mechanistic 4-Day Toxicity Study in the Male Mouse (Pharmacokinetic Investigations of the Clearance of Interavenously Administered 1251 Thyroxine), DACO: 4.8
1764323	2009, Fluopyram (AE C656948) Responses to the February 23, 2009 question form Germany BfR on metabolism and toxicology, DACO: 4.8,6.4
1764325	2009, Regulatory Position Paper Fluopyram: Response to PMRA on the in-house background incidence of "gall bladder absent" in the New Zealand White Rabbit fetus, DACO: 4.8
1764326	2009, Fluopyram (AE C656948) Weight of evidence evaluation of thyriod carcinogenesis in mice and liver carcinogenesis in rats using the IPCS mode of action framework, DACO: 4.8
1764327	2009, Hexyl Cinnamaldehyde (HCA), Potassium Dichromate (PDC) and Formaldehyde (FRM) Evaluation of Potential Dermal Sensitization in the Local Lymph Node Assay in the Mouse, DACO: 4.2.9
1764328	2008, AE 1801486: Evaluation of potential dermal sensitization in the local lymph node assay in the mouse, DACO: 4.2.9
1599345	2007, AE C656948: Comparative in vitro dermal absorption study in SC 500 formulation using human and rat skin, November 30, 2007. SA 07123, M-295237-01, ASB2008-5188, DACO 5.8, IIIA 7.6.2
1674473	2008, Fluopyram (AE C656948) SC 500 in vivo dermal absorption study in the male rat. August 26, 2008. SA 08082, Lynx-PSI N° TXGMP025, ASB2008-8225, DACO 5.8, IIIA 7.6.1
1599584	2008, AE C656948 500 SC - Magnitude of the residue on grape processed commodities, DACO: 7.4.5,IIA 6.5.3
1599586	2008, AE C656948 500 SC - Magnitude of the residue on small fruit vine climbing subgroup 13F, except fuzzy kiwifruit, DACO: 7.4.1,7.4.2,7.4.6,IIA 6.3.1
1599587	2008, AE C656948 500 SC: Magnitude of the residue in/on low growing berry (crop subgroup 13G), DACO: 7.4.1,7.4.2,7.4.6,IIA 6.3.2
1599619	2008, An analytical method for the determination of residues of AE C656948 in crop matrices using LC/MS/MS, DACO: 7.2.1,7.2.4,7.2.5,8.2.2.4,IIA 4.3
1599621	2008, Analytical method 00984 for the determination of residues of AE C656948 and its metabolites (AE F148815, AE C657188, BCS-AA10139, BCS-AA10065 and AE 1344122) and tebuconazole in/on plant material by HPLC-MS/MS, DACO: 7.2.1,7.2.4,7.2.5,8.2.2.4,IIA 4

1599624 2008, Analytical method 01061 for the determination of residues of fluopyram (AE C656948) and its metabolites AE F148815, BCS AA 10627 and BCS AA 10650 in/on animal tissues, milk and eggs by HPLC-MS/MS, DACO: 7.2.1,7.2.4,8.2.2.4,IIA 4.3 1599626 2008, Analytical method 01079 for the determination of residues of fluopyram (AE C656948) and AE F148815 in/on animal tissues, eggs and milk by HPLC-MS/MS, DACO: 7.2.1,7.2.2,7.2.4,8.2.2.4,IIA 4.3 1599645 2008, Determination of the residues of AE C656948 in/on grape (bunch of grapes) and bunch of grapes for wine proc. and the processed fractions (juice; raw juice; washings; pomace, dried; pomace, wet; berry, washed; retentate; pomace, grape; must; wine at 1st taste test; wine) after low-volume spraying of AE C656948 (500 SC) in the field in Southern France, DACO 7.4.5, IIA 6.5.3 1599646 2008, Determination of the residues of AE C656948 in/on grape (bunch of grapes) and bunch of grapes for wine proc. and the processed fractions (juice; raw juice; washings; pomace, dried; pomace, wet; berry, washed; retentate; pomace, grape; must; wine at 1st taste test; wine) after low-volume spraying of AE C656948 (500 SC) in the field in Southern France, DACO 7.4.5, IIA 6.5.3 1599658 2008, Determination of the residues of AE C656948 in/on strawberry fruit and the processed fractions (fruit, washed; preserve; washings; jam) after spraying of AE C656948 (500 SC) in the field in Northern France and Belgium, DACO: 7.4.5,IIA 6.5.3 1599659 2008, Determination of the residues of AE C656948 in/on strawberry fruit and the processed fractions (fruit, washed; preserve; washings; jam) after spraying of AE C656948 (500 SC) in the field in Southern France and Spain, DACO: 7.4.5, IIA 6.5.3 1599660 2008, Determination of the residues of AE C656948 in/on table grape (bunch of grapes) and the processed fractions (raisin; raisin waste; washings) after spraying of AE C656948 (500 SC) in the field in Spain, Portugal, Italy and Greece, DACO: 7.4.5,IIA 6 1599737 2008, Extraction efficiency testing of the residue analytical method 00984/M001 for the determination of AE C656948 residues in grapes using aged radioactive residues, DACO: 7.2.1,7.2.4,IIA 4.3 2008, Fluopyram: Feeding Study Laying Hens (Gallus gallus domesticus), DACO: 1599760 7.5.7.6.IIA 6.4.1 1599761 2008, Fluopyram: Feeding study with dairy cows, DACO: 7.5,7.6,IIA 6.4.2 1599769 2008, Independent laboratory validation of the analytical method 01079 for the determination of residues of fluopyram (AE C656948) and AE F148815 in/on animal tissues, eggs and milk by HPLC-MS/MS, DACO: 7.2.1,7.2.2,7.2.3,7.2.4,IIA 4.3 1599793 2008, Modification M001 of the analytical method 00984 for the determination of residues of AE C656948 and its metabolites (AE F148815, AE C657188 and BCS AA10139) and tebuconazole in/on plant material by LC-MS/MS, DACO: 7.2.1,7.2.4,7.2.5,8.2.2.4,IIA 4.3 1599801 2008, Phase report: 6 months stability in orange of study 07-02 - Storage stability of residues of AE C656948 and its metabolites (AE F148815, AE C657188 and BCS-AA10139) in orange during deep freeze storage for up to 24 months, DACO: 7.3, IIA 6.1.1

1599821 2008, Storage stability of residues of AE C656948 and its metabolites (AE F148815, AE C657188, BCS-AA10139 and BCS-AA10065) in plants during deep freeze storage for up to 24 months, DACO: 7.3,IIA 6.1.1 2009, Storage stability of residues of AE C656948 and its metabolites (AE 1784472 F148815, AE C657188, BCS-AA10139 and BCS-AA10065) in plants during deep freeze storage for up to 36 months - Progress interim report, DACO: 7.3 2009, Storage stability of residues of AE C656948 and its metabolites (AE 1804905 F148815, AE C657188 and BCS-AA10139) in orange during deep freeze storage for up to 36 months - Progress interim report (Phase Report: 24 months stability in orange of study 07-02), DACO 7.3 2008, Degradation of [phenyl-UL-14C] and [pyridyl-2,6-14C]AE C656948 by 1599640 plant suspension cell cultures, DACO: 6.3, IIA 6.2.1 1599779 2007, Metabolism of [phenyl-UL-14C]AE C656948 in beans after spray application, DACO: 6.3, IIA 6.2.1 2008, Metabolism of [phenyl-UL-14C]AE C656948 in confined rotational crops, 1599780 DACO: 6.3,7.4.4,IIA 6.2.1,IIA 6.6.2 2007, Metabolism of [phenyl-UL-14C]AE C656948 in potatoes, DACO: 6.3,IIA 1599781 6.2.1 1599782 2008. Metabolism of [phenyl-UL-14C]AE C656948 in red pepper after drip application, DACO: 6.3, IIA 6.2.1 1599785 2007, Metabolism of [phenyl-UL14C]AE C656948 in grapes after spray application, DACO: 6.3, IIA 6.2.1 1599786 2007. Metabolism of [pyridyl-2,6-14C]AE C656948 in grapes after spray application, DACO: 6.3, IIA 6.2.1 1599787 2008, Metabolism of [pyridyl-2,6-14C]AE C656948 in beans after spray application, DACO: 6.3, IIA 6.2.1 1599788 2008, Metabolism of [pyridyl-2,6-14C]AE C656948 in confined rotational crops, DACO: 6.3,7.4.4,IIA 6.2.1,IIA 6.6.2 1599789 2007, Metabolism of [pyridyl-2,6-14C]AE C656948 in potatoes, DACO: 6.3,IIA 6.2.1 1599790 2008, Metabolism of [pyridyl-2,6-14C]AE C656948 in red pepper after drip application, DACO: 6.3, IIA 6.2.1 2008, Metabolism of [phenyl-UL-14C]AE C656948 in the lactating goat, DACO: 1599783 6.2, IIA 6.2.3 1599784 2008, Metabolism of [phenyl-UL-14C]AE C656948 in the laying hen, DACO: 6.2, IIA 6.2.2 1599791 2008, Metabolism of [pyridyl-2,6-14C]AE C656948 in the lactating goat, DACO: 6.2, IIA 6.2.3 1599792 2008, Metabolism of [pyridyl-2,6-14C]AE C656948 in the laying hen, DACO: 6.2, HA 6.2.2 2010, Storage stability of residues of AE C656948 and its metabolites (AE 1983731 F148815, AE C657188, BCS-AA10139 and BCS-AA10065) in plants during deep freeze storage for up to 36 months, DACO: 7.3 2010, Storage stability of residues of AE C656948 and its metabolites (AE 1983732 F148815, AE C657188 and BCS-AA10139) in orange during deep freeze storage for up to 36 months, DACO: 7.3 1654363 2008, AE C656948 500 SC - Magnitude of the residue in/on crop tuberous and corm vegetables (crop subgroup 1C), DACO: IIA 6.3.8

1654364	2008, AE C656948 500 SC - Magnitude of the residue in/on sugar beets and leaves of root and tuber vegetables (crop group 2), DACO: IIA 6.3.10,IIA 6.3.9
1654372	2008, AE C656948 500 SC - Magnitude of the residue in/on peanut processed commodities, DACO: 7.4.5,IIA 6.5.3
1654373	2008, AE C656948 500 SC - Magnitude of the residue in/on field corn processed commodities and aspirated grain fractions, DACO: 7.4.5,IIA 6.5.3
1654374	2008, AE C656948 500 SC - Magnitude of the residue in/on wheat processed commodities and aspirated grain fractions, DACO: 7.4.5,IIA 6.5.3
1654375	2008, AE C656948 500 SC - Magnitude of the residue in/on soybean processed commodities and aspirated grain fractions, DACO: 7.4.5,IIA 6.5.3
1654376	2008, AE C656948 500 SC - Magnitude of the residue in/on cotton processed commodities, DACO: 7.4.5,IIA 6.5.3
1654378	2008, AE C656948 500 SC: Magnitude of the residue in/on canola processed commodities, DACO: 7.4.5,IIA 6.5.3
1654379	2008, AE C656948 500 SC - Magnitude of the residue in/on sugar beet processed commodities, DACO: 7.4.5,IIA 6.5.3
1654380	2008, AE C656948 500 SC - Magnitude of the residue in/on potato processed commodities, DACO: 7.4.5,IIA 6.5.3
1654383	2007, AE C656948 500 SC - Magnitude of the residue on apple processed commodities, DACO: 7.4.5,IIA 6.5.3
1654391	2007, Determination of the residues of AE C656948 in/on winter rape and summer rape and the processed fractions (oil, refined; oil, screwpressed; crude oil;) after spraying of AE C656948 (500 SC) in the field in Southern France and Italy,
1654393	DACO: 7.4 2007, Determination of the residues of AE C656948 in/on apple fruit and the processed fractions (fruit, washed; raw sauce; sauce; washings; strain rest; juice; pomace, wet; pomace, dried; raw juice; fruit, dried, peel rest; fruit, peeled) after spraying of AE C656948 (500 SC) in the field in Southern France and Italy, DACO 7.4.5, IIA 6.5.3
1654394	2007, Determination of the residues of AE C656948 in/on apple fruit and the processed fractions (fruit, washed; raw sauce; sauce; washings; strain rest; juice; pomace, wet; pomace, dried; raw juice; fruit, dried, peel rest; fruit, peeled) after spraying of AE C656948 (500 SC) in the field in Belgium and the United Kingdom, DACO 7.4.5, IIA 6.5.3
1654395	2007, Determination of the residues of AE C656948 in/on winter rape seed and the processed fractions (oil, refined; oil, screwpressed; crude oil; extracted meal; oil, solv. extracted; pomace) after spraying of AE C656948 (500 SC) in the field in, Germany, DACO 7.4.5, IIA 6.5.3
1654399	2008, AE C656948 500 SC - Magnitude of the residue in cotton (rotational crop tolerance), DACO: 7.4.4,IIA 6.6.3
1654400	2008, AE C656948 500 SC - Magnitude of the residue in field rotational crops (240-day plant back interval), DACO: 7.4.4,IIA 6.6.3
1654401	2008, AE C656948 500 SC - Magnitude of the residue in alfalfa (rotational crop tolerance), DACO: 7.4.4,IIA 6.6.3
1661215	2008, AE C656948 500 SC - Magnitude of the residue in/on dried, shelled peas and beans and the foliage of legume vegetables (crop subgroups 6C and 7A), DACO: 7.4.1,7.4.2,7.4.6,IIA 6.3.1

1661216	2008, AE C656948 500 SC - Magnitude of the residue in/on soybean, DACO: 7.4.1,7.4.2,7.4.6,IIA 6.3.1
1661219	2008, AE C656948 500 SC - Magnitude of the residue in/on cucurbit vegetables (crop group 9), DACO: 7.4.1,7.4.2,7.4.6,IIA 6.3.1
1661222	2008, AE C656948 500 SC - Magnitude of the residue in/on pome fruit (CG 11), DACO: 7.4.1,7.4.2,7.4.6,IIA 6.3.1
1661231	2008, AE C656948 500 SC - Magnitude of the residue on stone fruit, DACO:
1661238	7.4.1,7.4.2,7.4.6,IIA 6.3.1 2008, AE C656948 500 SC - Magnitude of the residue on tree nuts, DACO:
1661247	7.4.1,7.4.2,7.4.6,IIA 6.3.1 2008, AE C656948 500 SC - Magnitude of the residue in/on wheat and sorghum
1661248	(as part of crop groups 15 and 16, expect rice), DACO: 7.4.1,7.4.2,7.4.6,IIA 6.3.1 2008, AE C656948 500 SC - Magnitude of the residue in/on field corn and sweet corn (as part of crop groups 15 and 16, expect rice), DACO: 7.4.1,7.4.2,7.4.6,IIA 6.3.1
1661252	2008, AE C656948 500 SC - Magnitude of the residue in/on peanuts, DACO: 7.4.1,7.4.2,7.4.6,IIA 6.3.1
1661254	2008, AE C656948 500 SC - Magnitude of the residue in/on canola, DACO: 7.4.1,7.4.2,7.4.6,IIA 6.3.1
1661260	2008, AE C656948 500 SC - Magnitude of the residue in/on bananas (import
1661266	tolerance), DACO: 7.4.1,7.4.2,7.4.6,IIA 6.3.1 2008, AE C656948 500 SC - Magnitude of the residue in/on crop tuberous and
1661267	corm vegetables (crop subgroup 1C), DACO: IIA 6.3.8 2008, AE C656948 500 SC - Magnitude of the residue in/on sugar beets and
1661275	leaves of root and tuber vegetables (crop group 2), DACO: IIA 6.3.10,IIA 6.3.9 2008, AE C656948 500 SC - Magnitude of the residue in/on peanut processed
1661276	commodities, DACO: 7.4.5,IIA 6.5.3 2008, AE C656948 500 SC - Magnitude of the residue in/on field corn processed commodities and aspirated grain fractions, DACO: 7.4.5,IIA 6.5.3
1661280	2008, AE C656948 500 SC - Magnitude of the residue in/on wheat processed commodities and aspirated grain fractions, DACO: 7.4.5,IIA 6.5.3
1661282	2008, AE C656948 500 SC - Magnitude of the residue in/on soybean processed commodities and aspirated grain fractions, DACO: 7.4.5,IIA 6.5.3
1661283	2008, AE C656948 500 SC - Magnitude of the residue in/on cotton processed commodities, DACO: 7.4.5,IIA 6.5.3
1661285	2008, AE C656948 500 SC: Magnitude of the residue in/on canola processed commodities, DACO: 7.4.5,IIA 6.5.3
1661286	2008, AE C656948 500 SC - Magnitude of the residue in/on sugar beet processed commodities, DACO: 7.4.5,IIA 6.5.3
1661287	2008, AE C656948 500 SC - Magnitude of the residue in/on potato processed commodities, DACO: 7.4.5,IIA 6.5.3
1661291	2007, AE C656948 500 SC - Magnitude of the residue on apple processed
1661299	commodities, DACO: 7.4.5,IIA 6.5.3 2008, AE C656948 500 SC - Magnitude of the residue in cotton (rotational crop
1661301	tolerance), DACO: 7.4.4,IIA 6.6.3 2008, AE C656948 500 SC - Magnitude of the residue in field rotational crops (240-day plant back interval), DACO: 7.4.4,IIA 6.6.3

2008, AE C656948 500 SC - Magnitude of the residue in alfalfa (rotational crop tolerance), DACO: 7.4.4,IIA 6.6.3
2010, Fluopyram Projected Percent Crop Treated, DACO: 10.7.2 CBI
2010, Fluopyram Projected Percent Crop Treated, DACO: 10.7.2
2010, Projections of Percent crop Treated with fluopyram products in Canada, DACO: 10.7.2 CBI
2008, AE C656948 500 SC - Magnitude of the residue in/on pome fruit (CG 11), DACO: 7.4.1,9.3.2,IIA 8.3.1.1

3.0 Environment

PMRA Document Number	Reference
1599766	2008, Independent laboratory validation of analytical method 01023 for the determination of residues of AE C656948 and its metabolites AE C656948-benzamide (AE F148815), AE C656948-7-hydroxy (BCS-AA-10065) and AE C656948-PCA in soil by HPLC-MS/MS on soil, DACO: 8.2.2.1,8.2.2.2,IIA 4.4,IIA 4.6
1599620	2006, Analytical method 00973 for the determination of residues of AE C656948 in soil by HPLC-MS/MS, DACO: 8.2.2.1,IIA 4.4
1599622	2007, Analytical method 01023 for the determination of residues of AE C656948 and its metabolites AE C656948-benzamide (AE F148815), AE C656948-7-hydroxy (BCS-AA-10065) and AE C656948-PCA in soil by HPLC-MS/MS, DACO: 8.2.2.1,IIA 4.4
1599625	2008, Analytical Method 01068 for the determination of residues of AE C656948 in soil by HPLC-MS/MS, DACO: 8.2.2.1,IIA 4.4
1599627	2008, Analytical method for the determination of residues of AE C656948 and its metabolites AE C656948-benzamide, AE C656948-7-hydroxy, and AE C656948-PCA in soil and sediment using LC/MS/MS, DACO: 8.2.2.2,IIA 4.6
1599623	2007, Analytical method 01051 for the determination of fluopyram (AE C656948) in drinking and surface water by HPLC-MS/MS, DACO: 8.2.2.3,IIA 4.5
1599642	2008, Determination of fluopyram (AE C656948) in water by LC/MS/MS, DACO: 8.2.2.3,IIA 4.5
1599767	2008, Independent laboratory validation of analytical method 01051 for the determination of fluopyram (AE C656948) in drinking and surface water by HPLC-MS/MS, DACO: 8.2.2.3,IIA 4.5
1599507	2007, [14C]-AE C656948: Aqueous hydrolysis at pH 4, 7 and 9, DACO: 8.2.3.2,IIA 2.9.1,IIA 7.5
1599609	2007, AE C656948: Determination of the quantum yield and assessment of the environmental half-life of the direct photodegradation in water, DACO: 8.2.3.3,8.2.3.3.2,IIA 2.9.3,IIA 2.9.4,IIA 7.6
1599510	2008, [Phenyl-UL-14C)AE C656948 and [pyridyl-2,6-14C]AE C656948: Phototransformation in natural water, DACO: 8.2.3.3,8.2.3.3.2,IIA 2.9.4,IIA 7.6
1599509 1599508	2007, [14C]-AE C656948: Soil photolysis, DACO: 8.2.3.3.1,IIA 7.1.3 2008, [14C]-AE C656948: Aqueous photolysis in buffer at pH 7, DACO: 8.2.3.3.2,IIA 2.9.2,IIA 7.6

1599608	2007, AE C656948: Calculation of the chemical lifetime in the troposphere, DACO: 8.2.3.3.3,IIA 2.10,IIA 7.10
1599516	2008, [Phenyl-UL-14C]AE C656948: Aerobic soil metabolism/degradation and time-dependent sorption in four soils, DACO: 8.2.3.4.2,8.2.4.2,IIA 7.1.1,IIA 7.2.1,IIA 7.2.3,IIA 7.4.1
1599527	2008, [Pyridyl-2,6-14C]AE C656948: Aerobic metabolism/degradation and time-dependent sorption in soils, DACO: 8.2.3.4.2,8.2.4.2,IIA 7.1.1,IIA 7.2.1,IIA 7.2.3,IIA 7.4.1
1599511	2008, [Phenyl-UL-14C] and [pyridyl-2,6-14C]AE C656948: Aerobic soil metabolism in two US soils, DACO: 8.2.3.4.2,IIA 7.1.1,IIA 7.2.1
1774640	2009, Fluopyram - Bayer CropScience Response to PMRA Comments Regarding Redox Potential (Eh) in Anaerobic Soil and Aquatic Studies, DACO: 8.2.3.4.4,8.2.3.5.5,8.2.3.5.6,IIA 7.1.2,IIA 7.8.2
1599512	2008, [Phenyl-UL-14C] and [pyridyl-2,6-14C]AE C656948: Anaerobic soil metabolism, DACO: 8.2.3.4.4,IIA 7.1.2,IIA 7.2.4
1599531	2007, [pyridyl-ring-UL-14C]-AE C656948 and [triflurobenzamide-ring-UL-14C]-AE C656948 - Aerobic aquatic metabolism, DACO: 8.2.3.5.4,8.2.3.6,IIA 7.8.3
1599506	2007, [14C-pheny-UL]AE C656948: Anaerobic aquatic metabolism, DACO: 8.2.3.5.5,8.2.3.5.6,IIA 7.8.2
1599528	2007, [pyridyl-2,6-14C]AE C656948: Anaerobic aquatic metabolism, DACO: 8.2.3.5.5,8.2.3.5.6,IIA 7.8.2
1599772	2008, Kinetic evaluation of the aerobic aquatic metabolism of fluopyram (AE C656948) in water/sediment systems using MatLab, DACO: 8.2.3.6,IIA 7.8.3
1599607	2007, AE C656948: Adsorption/desorption on five soils, DACO: 8.2.4.2, IIA 7.4.1
1599735	2008, Evaluation of the time-dependent sorption of fluopyram (AE C656948) based on laboratory batch equilibrium experiments in 8 soils, DACO: 8.2.4.2,IIA 7.4.1
1599520	2007, [Pyridine-2,6-14C] AE C656948-7-hydroxy: Adsorption/desorption on four EU soils, DACO: 8.2.4.2, IIA 7.4.2
1599751	2008, Fluopyram - Statement on the pyrolytic behaviour under controlled conditions and on the controlled incineration as a safe means of disposal - AE C656948, DACO: 8.4.1,IIA 3.8.1
1599652	2007, Determination of the residues of AE C656948 in/on soil after spraying of AE C656948 (250 SC) in the field in Germany, United Kingdom, Sweden, France Spain and Italy, DACO: 8.6,IIA 7.3.1
1599771	2008, Kinetic evaluation of field dissipation studies after application of fluopyram (AE C656948) in Europe according to FOCUS using KinGui, DACO: 8.6,IIA 7.3.1
1599497	2007, 1. Interim Report: Determination of the residues of AE C656948 in/on soil after spraying of AE C656948 (250 SC) in Germany and France, DACO: 8.6,IIA 7.3.3
1599606	2008, AE C656948: Acute toxicity to earthworms (<i>Eisenia fetida</i>) tested in artificial soil with 5 percent peat, DACO: 9.2.3.1,IIA 8.9.1
1599589	2008, AE C656948 SC 500: Effects on survival, growth and reproduction on the earthworm <i>Eisenia fetida</i> tested in artificial soil with 5 percent peat, DACO: 9.2.3.1,IIA 8.9.2
1599733	2007, Effects of AE C656948 (acute contact an oral) on honey bees (<i>Apis mellifera L.</i>) in the laboratory, DACO: 9.2.4.1,9.2.4.2,IIA 8.7.1,IIA 8.7.2

2008, Dose-response toxicity (LR50) of AE C656948 SC 500 to the predatory 1599727 mite Typhlodromus pyri (Scheuten) under laboratory conditions, DACO: 9.2.5, IIA 8812 2008, Dose-response toxicity (LR50) of AE C656948 SC 500 to the parasitic wasp 1599729 Aphidius rhopalosiphi (Destefani-Perez) under laboratory conditions, DACO: 9.2.6.IIA 8.8.1.1 2008. AE C656948 tech.: Determination of effects on nitrogen transformation in 1599599 soil, DACO: 9.2.8,9.2.9,IIA 8.10.1 1599593 2008. AE C656948 tech.: Determination of effects on carbon transformation in soil, DACO: 9.2.8.9.2.9, IIA 8.10.2 2007, Acute toxicity of AE C656948 (tech.) to the waterflea Daphnia magna in a 1599541 static laboratory test system, DACO: 9.3.2,IIA 8.3.1.1 1599770 2008, Influence of AE C656948 (tech.) on development and reproductive output of the waterflea Daphnia magna in a static renewal laboratory test system, DACO: 9.3.3.IIA 8.3.2.1 1599592 2008, AE C656948 SC 500A G: Influence on the reproduction of the collembola species Folsomia candida tested in artificial soil with 5 % peat, DACO: 9.3.4,9.6.6,9.9,IIA 8.16.1 2008, AE C656948 tech.: Determination of effects on growth of pure cultures of a 1599594 soil fungus, Agrocybe aegerita, on a soil-nutrient medium, DACO: 9.3.4,9.6.6,9.9,IIA 8.16.1 1599595 2008, AE C656948 tech.: Determination of effects on growth of pure cultures of a soil fungus, Cladorrhimum foecundissimum, on a soil-nutrient medium, DACO: 9.3.4,9.6.6,9.9,IIA 8.16.1 2008, AE C656948 tech.: Determination of effects on growth of pure cultures of a 1599596 soil fungus, Mucor circinelloides var. griseocyanus, on a soil-nutrient medium, DACO: 9.3.4,9.6.6,9.9,IIA 8.16.1 2008, AE C656948 tech.: Determination of effects on growth of pure cultures of a 1599597 soil fungus, Penicillium simplicissimum, on a soil-nutrient medium, DACO: 9.3.4,9.6.6,9.9,IIA 8.16.1 2008, AE C656948 tech.: Determination of effects on growth of pure cultures of a 1599598 soil fungus, Phytophthora nicotianae, on a soil-nutrient medium, DACO: 9.3.4,9.6.6,9.9,IIA 8.16.1 2008, Chronic dose-response toxicity (ER50) of AE C656948 SC 500 to the rove 1599634 beetle Aleochara bilineata GYLL. under extended laboratory conditions, DACO: 9.3.4,9.6.6,9.9,IIA 8.16.1 1599752 2008, Fluopyram SC 500: Influence on mortality and reproduction on the soil mite species Hypoaspis aculeifer tested in artificial soil with 5 % peat, DACO: 9.3.4,9.6.6,9.9.IIA 8.16.1 1599603 2007, AE C656948: A 96-hour flow-through acute toxicity test with the saltwater mysid (Americamysis bahia), DACO: 9.4.2,9.4.3,9.4.4,IIA 8.11.1 2006, AE C656948: A 96-hour shell deposition test with the eastern oyster 1599604 (Crassostrea virginica), DACO: 9.4.2,9.4.3,9.4.4,IIA 8.11.1 2008, Acute toxicity of AE C656948 (tech.) to fish (Oncorhynchus mykiss) under 1599539 static conditions, DACO: 9.5.2.1,9.5.2.3,IIA 8.2.1.1 1599537 2008. Acute toxicity of AE C656948 (tech.) to fish (Cyprimus carpio) under static conditions, DACO: 9.5.2.2,9.5.2.3,IIA 8.2.1.2

1599538	2008, Acute toxicity of AE C656948 (tech.) to fish (Lepomis macrochirus) under
1500543	static conditions, DACO: 9.5.2.2,9.5.2.3,IIA 8.2.1.2
1599543	2008, Acute toxicity of AE C656948 technical to the fathead minnow (<i>Pimephales promelas</i>) under static conditions, DACO: 9.5.2.2,9.5.2.3,IIA 8.2.1.2
1599544	2006, Acute toxicity of AE C656948 technical to the sheepshead minnow
	(Cyprinodon variegatus) under static conditions, DACO: 9.5.2.4,IIA 8.11.1
1599730	2007, Early-life stage toxicity of AE C656948 (tech.) to fish <i>Pimephales</i>
	promelas), DACO: 9.5.3.1,IIA 8.2.4
1599536	2008, Acute oral toxicity for bobwhite quail (Colimus virginianus) with AE
	C656948 techn. a.s., DACO: 9.6.2.1,9.6.2.2,9.6.2.3,IIA 8.1.1
1654429	2008, AE C656948 - Acute oral toxicity test (LD50) with the zebra finch
	(Taeniopygia guttata) following OECD draft guideline 223, DACO:
	9.6.2.1,9.6.2.2,9.6.2.3,IIA 8.1.1
1661315	2008, AE C656948 - Acute oral toxicity test (LD50) with the zebra finch
	(Taeniopygia guttata) following OECD draft guideline 223, DACO:
	9.6.2.1,9.6.2.2,9.6.2.3,IIA 8.1.1
1599554	2007, AE C656948 (tech. a.s.) - 5-day-dietary LC50 for bobwhite quail (Colimus
	virginiamus), DACO: 9.6.2.4,9.6.2.5,IIA 8.1.2
1599600	2007, AE C656948 techn. a.s.: 5-day-dietary LC50 mallard duck (Anas
	platyrhynchos), DACO: 9.6.2.5,9.6.2.6,IIA 8.1.3
1599605	2008, AE C656948: A reproduction study with the Northern bobwhite, DACO:
	9.6.3.1,9.6.3.2,9.6.3.3,IIA 8.1.4
1599731	2008, Effect of AE C656948 technical on reproduction of the mallard duck (Anas
	platyrhynchos), DACO: 9.6.3.1,9.6.3.2,9.6.3.3,IIA 8.1.4
1599732	2008, Effect of AE C656948 technical on reproduction of the northern bobwhite
	quail, DACO: 9.6.3.1,9.6.3.2,9.6.3.3,IIA 8.1.4
1599588	2008, AE C656948 SC 500: Effects on soil litter degradation, DACO:
	9.6.6,9.9,IIA 8.16.2
1599808	2008, Pseudokircheriella subcapitata growth inhibition test with fluopyram-
	lactame, DACO: 9.8.2,9.8.3,IIA 8.4
1599862	2007, Toxicity of AE C656948 technical to the 2007, freshwater diatom <i>Navicula</i>
	pelliculosa, DACO: 9.8.2,9.8.3,IIA 8.4
1599863	2007, Toxicity of AE C656948 technical to the blue-green algae <i>Anabaena flos-</i>
	aquae, DACO: 9.8.2,9.8.3,IIA 8.4
1599864	2007, Toxicity of AE C656948 technical to the green alga <i>Pseudokirchneriella</i>
	subcapitata, DACO: 9.8.2,9.8.3,IIA 8.4
1599865	2007, Toxicity of AE C656948 technical to the saltwater diatom <i>Skeletonema</i>
	costatum, DACO: 9.8.3,IIA 8.11.1
1599590	2008, AE C656948 SC 500A G - Effect on the vegetative vigour of ten species of
	non-target terrestrial plants (Tier 1), DACO: 9.8.4,IIA 8.12
1599591	2008, AE C656948 SC 500A G effect on seedling emergence and seedling growth
	test of ten species of non-target terrestrial plants (Tier 1 and 2), DACO: 9.8.4,IIA
	8.12
1599734	2008, Evaluation of the pre-emergence (PPI) biological activity of AE C656948
	SC 500, DACO: 9.8.4,IIA 8.12
1599773	2007, Lemna gibba G3 - Growth inhibition test with AE C656948 under static
	conditions, DACO: 9.8.5,IIA 8.6
1599602	2008, AE C656948- Toxicity to bacteria, DACO: 9.9,IIA 8.15

2008, AEC656948 - Toxicity to marine amphipods (*Leptocheirus plumulosus*) during a 10-day sediment exposure, DACO: 9.9,IIA 8.5.1
 2008, AEC656948 - Life-cycle toxicity test exposing midges (*Chironomus tentans*) to a test substance applied to sediment under static-renewal conditions following EPA test methods, DACO:
 2008, AEC656948 - Toxicity to estuarine amphipods (*Leptocheirus plumulosus*) during a 28-day sediment exposure, DACO: 9.9,IIA 8.5.2
 2008, *Chironomus riparius* 28-day chronic toxicity test with fluopyram (tech.) in a water-sediment system using spiked water, DACO: 9.9,IIA 8.5.2

4.0 Value

PMRA Document	Reference
Number	
1599332	2008. Fluopyram 500 SC Fungicide (500 g a.i./L fluopyram) for control of Botrytis bunch rot in grape, botrytis grey mould in strawberry and tomato, and
	Alternaria solani in tomato, DACO: 10.2.3.3, 10.2.3.4, 10.3.2, 10.4, 10.5.1,
	10.5.2, 10.5.4, IIIA 6.1.2, IIIA 6.1.3, IIIA 6.2.1, IIIA 6.3, IIIA 6.4.1, IIIA 6.4.2, IIIA 6.4.3, 322pp.
1670787	2008. Fluopyram/Prothioconazole Fungicide for Control of Ascochyta Blight of Lentil and Chickpea, Mycosphaerella Blight of Dried Shelled Pea, and White
	Mold of Dried Shelled Bean and Pea. DACO: 10.2.3.3, 10.2.3.4, 10.3.2, 10.4, 10.5.1, 10.5.2, 10.5.4. 272pp.
1670080	2008. Fluopyram + pyrimethanil 500 SC fungicide (125g a.i./L fluopyram + 375g a.i./L pyrimethanil) for control of listed diseases in grapes and small berries, bulbvegetables, tomatoes, and pome fruit. DACO: 10.2.3.3, 10.2.3.4, 10.3.2, 10.4, 10.5.1, 10.5.2, 10.5.4. 420pp.
1674457	2008. Fluopyram 500 SC Fungicide for control of listed diseases in horticulture and field crops, DACO: 10.2.3.3, 10.2.3.4, 10.3.2, 10.4, 10.5.1, 10.5.2, 10.5.4.
2046958	851pp. 2011. Cover Letter for Fluopyram Clarification request Sub No 2008-4863 efficacy data to add drip irrigation strawberries. DACO: 0.8. 2pp.
2046960	2011. Efficacy data. DACO: 10.5. 5pp.
2046961	2011. Efficacy data. DACO: 10.5. 5pp.
2046963	2011. Efficacy data. DACO: 10.5. 7pp.